Steered Molecular Dynamics Simulations of Coumarin2 5Z/5E Pulling Reveal Different Interaction Profiles for Four Human Cytosolic Carbonic Anhydrases

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Abstract: Carbonic anhydrase (CA) is an important enzyme, which has multiple isoforms each performing different functions in humans. Characterizing the interactions of carbonic anhydrase isoforms with ligands is a difficult but important task. In recent years, steered molecular dynamics (SMD) has been used successfully to characterize ligand binding and unbinding events. In this study, constant velocity SMD simulations were performed on four cytosolic human carbonic anhydrases (carbonic anhydrase I, II, III and VII) using a ligand called Coumarin2 5Z/5E to understand better the nature of interactions between this ligand and the various enzyme isoforms. The influence of force constant and pulling velocity on SMD simulations were investigated. When the force constant and the pulling velocity were changed, no significant effect on interaction profiles was observed. Finally, hydrogen bond interactions, Histidine 64 positions and the differences between force-distance profiles for all isozymes were analyzed. These results demonstrate that human cytosolic carbonic anhydrases I, III, and VII show similar Coumarin2 5Z/5E unbinding patterns with different rupture forces, while carbonic anhydrase II has a distinct interaction profile.

Keywords: Steered molecular dynamics, Coumarins, Rupture force, Protein-ligand interaction, Human cytosolic carbonic anhydrases

1. Introduction

Carbonic anhydrases (CAs) are zinc metalloenzymes that play a vital role in various organisms, including humans. This vital role includes many physiological, as well as pathological processes, including lipogenesis, gluconeogenesis, ureagenesis, and tumorigenesis [1, 2]. There are 16 known isozymes of alpha-CAs which are found in different subcellular locations [1]. Five of them, namely CA I, II, III, VII and XII, function in the cytosol. Human CA I, II, III and VII share approximately 35% sequence identity and have
The zinc interacting histidines, namely His 64, 94, 96 and 119 (CA II numbering), are conserved across all four of these isoforms. However, small changes in sequence may cause striking differences in enzymatic activity.

Since carbonic anhydrase inhibitors are used as drugs to treat diseases like glaucoma, epilepsy, cancer, and others, finding selective inhibitors targeting a specific CA isoform is an important goal. A number of sulfonamide- and sulfamate-based CA inhibitors have been developed over time [3, 4]. Most of these inhibitors interact with zinc, and they do not form a covalent bond with CAs [5]. Therefore, they constitute a new class of CA inhibitors with a different mechanism of inhibition. It will be of tremendous benefit to understand, at the molecular level, the precise mechanisms by which Coumarin interacts with CAs, leading to inhibition of enzyme activity. Knowledge of Coumarin's enzyme specificity and a description of its interaction with specific CA amino acid residues could help the development of novel drugs, with better specificity, and less side effects, for the treatment of diseases like glaucoma, epilepsy and cancer [6, 7].

The interactions of three Coumarin derivatives (Coumarin1, Coumarin2, and Coumarin3) with various CA isoforms have been reported previously [5]. In that study, the hydrolysis product of Coumarin2 (referred to as Coumarin2 5Z/5E) was shown to have a greater binding affinity for CA I and CA II than Coumarin2 itself. Unfortunately, that study only reported the binding affinity of Coumarin2 5Z/5E for CA I and CA II. Therefore, it is still unknown how Coumarin2 5Z/5E interacts with other cytosolic CAs like CA III and CA VII. Likewise, the molecular details of its interactions with CAs and the precise residues involved in binding remain to be elucidated. For these reasons, the author investigated how Coumarin2 5Z/5E interacts with four human cytosolic CAs (namely CAs I, II, III and VII). Although there are numerous atomic level studies on carbonic anhydrases, most of these have concentrated on the interactions with zinc and the well-described proton transfer mechanism [11-15]. Instead here, the author have focused on describing the atomic interactions of Coumarin2 5Z/5E with four human cytosolic CAs using SMD method.

SMD simulations are similar to single molecule Atomic Force Microscopy (AFM) experiments. These types of experiments can shed light on the interaction of ligands with proteins [16-18]. Atomic scale data of rupture forces, specific residue-ligand interactions, binding/unbinding pathways, ligand recognition residues, as well as the calculated potential of mean force, can all be derived as a result of SMD simulations [19-22]. Due to the tremendous wealth of information it can provide, SMD was used to characterize the interactions of Coumarin2 5Z/5E with four human cytosolic CAs.

In this study, answers of following three questions were sought: Can rupture force be used to quantify the degree of interaction of Coumarin2 5Z/5E with CAs? Is there a possible long-range effect of Coumarin2 5Z/5E on the zinc-interacting residue, Histidine 64, a residue known to be involved in proton transport? Can the residues that play an important role in the binding/unbinding of Coumarin2 5Z/5E be identified?

2. Material and Method

The following Protein Databank entries were used for simulations as initial structures: 1HCB for CA I, 3FBE for CA II, 1Z93 for CA III, and 3MDZ for CA VII [5, 23, 24]. Water molecules in the crystal structures were retained. The systems were solvated with a 10 Å solvent buffer to produce triclinic boxes. Na⁺ and Cl⁻ counter ions were added at 0.15 M to make the systems neutral. The NAMD program with Charmm27
force field was used to perform SMD simulations [25-27]. The system compositions for all simulations are given in Table S1.

A previous automated procedure was adopted to determine the atom types of Coumarin2 5Z/5E in the framework of the Charmm General Force Field (CGenFF) 0000 [28]. After automated atom type assignment, Coumarin2 5Z/5E was parameterized using the Force Field Tool Kit (FFTK) implemented in VMD [29]. The Gaussian 09 program was used for quantum mechanical calculations in the parameterization procedure [30].

A thorough minimization of 50000 steps was performed after system construction. The systems were heated to 300 K in 0.001 K increments and one femtosecond time step was used for all of the simulations. A two-nanosecond equilibration period, under constant pressure (1 atm.) and temperature (300 K), followed the heating procedure. The Langevin barostat and thermostat were used for pressure and temperature control [31]. Particle Mesh Ewald (PME) method, with a grid spacing of 1.0 Å, was used to model electrostatic interactions [32] with the long-range force cutoff distance set to 12.5 Å. The average Cα Root Mean Square Deviations (RMSDs) during equilibration were 0.756 Å, 0.844 Å, 0.851 Å and 0.973 Å for the CA I, II, III and VII crystal structures, respectively. Even though there was not any restriction during equilibration, the RMSD was small and this indicated that the enzymes did not undergo a conformational change during equilibration. Hydrophobic and hydrogen bond interactions of the final frames of the equilibration procedure were plotted using LigPlot+ and the data is shown in Supplementary Information Figure S3 [33].

Similar to AFM experiments, SMD simulations are also mostly single molecule studies. For this reason, a periodic boundary condition was not used during production runs. In production runs, the ligand was removed from the active site cavity for 2.5 nanoseconds along an approximately linear path with a constant velocity of 0.04 Å/ps. The production run time was doubled when the pulling velocity was reduced to 0.02 Å/ps. VMD program and various custom-made Python scripts were used for analyses [9]. The amino- and carboxy-terminals of all the CAs were fixed to prevent protein drift during ligand pulling and this remained the only applied constraint.

3. Results

3.1. Effect of different force constants on the force-time profile

Initially, the author wished to explore what effect changing the force constant would have on Force-Time profiles. In order to examine this, CA II was tested with both k=500 pN/Å and k=1000 pN/Å as the spring force constants. The Xmgrace program, using a running average of 25 frames, was used to remove noise from all force related curves (Figure 2). The results show that the Force-Time profiles were relatively insensitive to spring force constant change. However, the stiffer spring force constant (k=1000 pN/Å) was selected due to the recommendations in previous studies [34, 35].

Figure 2. Force-Time profiles for k=500 pN/Å (black) and k=1000 pN/Å (blue) force constants.

3.2. Force-distance profiles

Force in the pulling direction was plotted against the distance of the ligand from its initial position for ten trajectories with velocity v=0.04 Å/ps (Figure 3). A peak around 50 Å was found to be present in all of the plots. This peak corresponds to complete dissociation of the ligand from the protein and the solvation shells around it. Snapshots of the first trajectories, with approximately a 50 Å distance between the ligand and the individual CA isozymes, are shown in Figure S4. In a Force-Distance plot, the highest value of force is called the rupture force. The rupture force did not vary significantly between CA I and CA II. On the other hand, in CA III it was found to vary from approximately 523 pN (Trajectory 10) to 1153 pN (Trajectory 2) while the lowest was 312 pN (Trajectory 1). The force becomes approximately zero above a protein-ligand distance of 60 Å for all isozymes. A sharp peak around 20 Å was also common in all cytosolic CAs with the exception of CA II.

The selection of a reasonable velocity in constant velocity pulling simulations is an important consideration. Previous studies have shown that varying the pulling velocity does not significantly influence SMD results [20, 36]. In the majority of cases, reducing the pulling velocity results only in smaller force values. Indeed, qualitatively similar results were observed in this study as well. However, repeating SMD simulations with various velocities to ensure qualitative similarity of the results is a common practice [16, 36]. For this reason, all of the simulations were repeated with v=0.02 Å/ps velocity. The data of Force-Distance profiles is given in Figure 4.
As expected, and discussed above, the force decreased slightly in the slower simulations for almost all CA trajectories. Similar to the 0.04 Å/ps simulations, there was significantly more variation in force in the CA III simulations. Despite these differences, in general the shapes of all SMD simulations were similar regardless of the pulling velocity. As a result, the insensitivity of the data to the pulling velocity shows that the data is robust.
Figure 5. Average forces (lines) and their standard deviations (shades) for all simulations. A) Velocity=0.04 Å/ps simulations B) Velocity=0.02 Å/ps simulations.

The average forces and their standard deviations were plotted with respect to simulation time for all systems and the data is presented in Figure 5. The data clearly show that, although simulation times vary, the average force profiles are preserved for both the slow and the fast simulations. The average rupture force for CA III is about 700 pN at 0.04 Å/ps and it decreases to 600 pN when the pulling velocity is reduced to 0.02 Å/ps. The average rupture force of CA I is 500 pN for the faster simulation and it decreases to about 100 pN in the slower simulation. For CA VII, the average rupture force is 400 pN at 0.04 Å/ps pulling velocity and it falls to approximately 300 pN when the pulling velocity is reduced to 0.02 Å/ps. Interestingly, CA II has a distinct profile when compared with the other three CAs. The height of the force peak was not as sensitive to pulling velocity as was observed for the other CAs. It should also be noted that standard deviations are higher in the simulations for CA III and CA VII compared to CA I and II. The unprocessed Force-Time curves are presented in Supplementary Information Figure S1 and S2.

3.3. Position of histidine 64 (His 64)

His 64 (based on CA II residue numbering) plays an important role in the proton transfer mechanism of many CAs, and many sulfonamide- and sulfamate-based CA inhibitors interact with this residue. Because of its importance, His 64 positions in CA II have been the subject of computational and experimental studies [11, 15]. Based on the distance of ND1 atom of His 64 residue and Zn atom, His 64 has two conformations: namely “in” and “out” conformations. The author wondered whether Coumarin2 5Z/5E could cause any conformational change in the His 64 residue during the unbinding process (Figure 6). Since CA III does not have a corresponding Histidine (see Figure 1-C sequence alignment), it was excluded from this analysis. The data show that CA I has the greatest ND1-Zn distance. It turns to the “in” conformation in only two trajectories (Trajectory 4 and 6). Since this conformational change occurs before rupture of the ligand from the protein, it is not correlated with ligand unbinding. His64 in CA II demonstrate “out” conformations at approximately 8 Å distance and frequent jumps to the “in” conformation are observed. In contrast, although CA VII “out” conformations have distances similar to CA II, similar jumps to an “in” conformation were not observed. Based on these observations, which do not demonstrate a concerted His64 conformational change with Coumarin2 5Z/5E unbinding, the conclusion is that Coumarins interact with CAs in a manner distinct from sulfonamide- and sulfamate-based inhibitors.

3.4. Hydrogen bonds between Coumarin2 5Z/5E and CA isozymes

Hydrogen bonds formed between Coumarin2 5Z/5E and all of the CA isozymes were investigated during the simulation (Figure 7). A consistent hydrogen bond between Coumarin2 5Z/5E and Lys 58 of CA I was observed in the first half nanosecond. A short-lived hydrogen bond between the ligand and His 4 was noted for CA II; this bond had the shortest lifetime compared to other hydrogen bonds noted. Coumarin2 5Z/5E also formed a hydrogen bond with Arg 67 of CA III and, in contrast with the CA II His 4 bond, this hydrogen bond was the most stable of the three hydrogen bonds Coumarin2 5Z/5E formed with CA III. In fact, the lifetime of this hydrogen bond interaction extended up to 750 ps and it was the most persistent hydrogen bond noted among the four isozyme simulations. Lys 175 of CA VII also formed a hydrogen bond with Coumarin2 5Z/5E and the lifetime of this bond was comparable with the hydrogen bond of Arg 67 in CA III. However, CA VII Lys 175 formed only one bond with Coumarin2 5Z/5E. These data are perhaps not surprising since hydrogen bonding residues are not conserved across the different isozymes (see Figure 1-C). Additionally, one can clearly see that the difference between the hydrogen bonding patterns of Coumarin2 5Z/5E across isozyme may explain why there is a peak around 20 Å in CA I, III, VII and why this peak is not observed in CA II.
4. Discussion and Conclusion

In this study, 40 ligand-unbinding SMD simulations for four human cytosolic CAs were performed. Each enzyme showed a different pattern of interactions with the ligand Coumarin2 5Z/5E. The order of rupture force is CA III > CA I > CA VII > CA II. Previously published data has shown that Coumarin2 5Z/5E has a lower binding affinity for CA I than CA II [5]. These SMD simulations also confirm this fact qualitatively. Correlation between rupture forces and experimental binding affinities have been studied [22] and have shown that a higher rupture force is correlated with a lower binding affinity. Based on this correlation, one can predict that CA III will have the lowest binding affinity and CA II will have the highest binding affinity for Coumarin2 5Z/5E. Compared to the other CAs, CA II demonstrates the least ligand interactions. Long range interactions such as electrostatic and Van der Waals forces are predominant in the interactions of CA II with Coumarin2 5Z/5E. In contrast, CA III interacts strongly with Coumarin2 5Z/5E whereas the interactions of ligand with CA VII and CA I are intermediate between CA III and CA II.

These results also highlight both the important residues involved in Coumarin2 5Z/5E interactions, as well as differences in residue interaction across isozymes. Lys 58 in CA I, His 4 in CA II, Arg 67 in CA III and Lys 175 in CA VII are all important residues involved in interactions with Coumarin2 5Z/5E.
Mutations of these residues will significantly disrupt Coumarin2 5Z/5E inhibition. This new information can now be used to aid in designing a new generation of carbonic anhydrase inhibitors that could lead to new drugs for the treatment of glaucoma, epilepsy, cancer and other diseases. The data in this paper also confirm that, since there is no concerted motion between His 64 conformations and Coumarin2 5Z/5E, the mechanism of Coumarin inhibition is not through a zinc interaction or a proton transfer mechanism.

In summary, the SMD simulation of cytosolic CAs has successfully led to predictions regarding the relative affinity and binding mode of Coumarin2 5Z/5E to these enzymes. Although SMD can’t give experimental unbinding energies, it is still a very useful computational method to characterize possible ligand-protein interactions as well as average rupture forces of ligands, which are known to be correlated with experimental binding affinities as shown previously [22].

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References


Appendices

Appendix A. Supplementary Information

This supplementary information contains 4 figures and 1 table. Table A1 is behind the figures, at the last page of this document.

Figure A1. Force-Time curves for 0.04 Å/ps simulations. A) CA I B) CA II C) CA III D) CA VII

Figure A2. Force-Time curves for 0.02 Å/ps simulations. A) CA I B) CA II C) CA III D) CA VII
Figure A3. Coumarin 5Z/SE interactions when production simulations started for A) CA I, B) CA II, C) CA III and D) CA VII

Figure A4. Snapshots of protein and ligand halfway through simulations. Distance unit is Å. A) CA I B) CA II C) CA III D) CA VII

Table A1. System compositions for all simulations

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