Accurate Modeling of Cation- π interactions in Enzymes: A Case Study on the CDPChO:phosphocholine Cytidylyltransferase Complex

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ABSTRACT:

Cation- π interactions are functionally relevant, strong secondary interactions that play versatile roles in a variety of chemical and biological systems. Therefore it is very important to be able to describe accurately and reliably these interactions. In this study we propose a methodology for the accurate modelling of cation- π interactions in proteins using QM/MM calculations. We developed a methodology for computing the many-body interaction energy terms and tested the effect of various factors on the accuracy of the binding energy. We found that once well-equilibrated structures were reached in the MD simulations, very similar results can be obtained for the various snapshots taken from the trajectory. The calculated interaction energies were only slightly influenced by electrostatic embedding of the point charges in the QM/MM calculations, and by QM/MM geometry optimization. The calculated molecular mechanics interaction energies were off by 50% for cation- π interactions. Instead, we suggest the calibration of force fields based on fragment-based QM-calculations on geometries obtained from MD simulations to yield reliable binding energies at reduced computational cost.

Dedicated to Professor Magdolna Hargittai on the occasion of her 70th birthday.

1. Introduction

Non-covalent interactions, such as hydrogen bonds and salt bridges, play a crucial role in all biological systems.[1] Among others, they contribute to the three dimensional structure of biomolecules, to ligand binding and to enzymatic catalysis. As such, *in silico* drug discovery and design methodologies depend to a large extent on proper description of these intermolecular forces. In the course of virtual screening, millions of small compounds are docked into the ligand binding site of receptors or enzymes and scoring functions are used to estimate the strength and likelihood of binding.[2]

Cation- π interaction is a relatively newly recognized secondary molecular interaction, whereby a positively charged molecule or molecular moiety interacts with an aromatic ring.[3] The strength of a cation- π interaction is significant; it is comparable to hydrogen bonds and salt bridges.[4] Several intermolecular forces, such charge-quadrupole, charge-dipole, charge-transfer and dispersion forces, contribute to the formation of cation- π interactions, but the most significant energy contribution is derived from electrostatic interactions.[5,6]

Cation- π interactions play a prominent role in a variety of fields, e.g. in structural biology, materials science, catalysis and organic synthesis.[4,7] Their relevance in biological systems where they contribute significantly to the stabilization of protein structure and ligand binding, has been extensively demonstrated.[8]

The evaluation of 593 high-resolution structures in Protein Data Bank showed that (1) from every 77 residues 1 participates in cation- π interaction (2) arginine is more likely to be involved in cation- π interactions than lysine (3) 25% of tryptophan residues form energetically significant cation- π interaction.[9] Web servers, such as CaPTURE,[9] and Protein Explorer[10] use distance-based criteria to find structurally significant cation- π interactions and use force field parameters fitted to *ab initio* calculations to predict the strength of the interaction. These methods are fast, but very specific and they can only be applied certain types of interactions.

In contrast to molecular mechanics, quantum mechanics based methods have the ability to accurately describe inter- and intramolecular interactions in any system, but these methods are too computationally demanding for screening purposes. Therefore in the course of drug discovery and design usually scoring functions, approximate mathematical expressions, are used to approximate the binding energy in order to accelerate the computational process. One of the most frequently applied approaches is to use force fields to calculate the interaction energy between the protein and the ligand. This method is fast and easily applied, however, it has serious limitations, e.g. neglecting electronic polarization, approximate handling of van der Waals interactions, and difficult or no treatment of solvation effects. Although the parameterization of force fields is usually done with great care, extensive validation of the reliability of the force field can only be achieved during extended applications and repeated evaluation. Ideally one would like to get excellent agreement between calculated and real (measurable) binding energies, or at least obtain the same trend. Achieving such a qualitative agreement is a great advantage, and shows the reliability of the method.[11] When experimental data are not available, the computed results can be compared to data calculated at a higher level, e.g. results of MM methods could be compared to *ab initio* or DFT data.

An important advantage of computed data is that they can reveal the contribution of protein side chains to the overall binding energy of the ligand, which is very difficult to obtain experimentally. Therefore in this work we 1) design a methodology to obtain accurate binding energies at low computational cost and 2) tackle the factors involved in ligand binding. One of the major issues considered in accurate modeling of protein-ligand interactions is protein flexibility. Therefore we investigate (1) how thermal motion influences the results by studying the binding energy along the MD trajectory. (2) the generally used approximation of the calculation of the binding energy as a sum of pair potentials by calculating the many-body interaction energy terms (3) the possible improvement of QM/MM calculations over MM calculations and the effect of electrostatic embedding (4) the effect of DFT functional and basis set.

In this work, we use the phosphocholine cytidylyltransferase enzyme (*Pf*CCT) from *Plasmodium Falciparum* causing malaria as a model system. This protein has been recently actively researched as a potentially new drug target against malaria.[12,13] Furthermore, detailed information exists on the role of the various active site side-chains on ligand binding affinity and enzyme activity, and detailed thermodynamic properties of point mutants, which will enable us to compare the computed results to experiments.[14]

*Pf*CCT functions as a homodimer (Fig. 1) and plays a key role in the membrane synthesis of the pathogen of malaria. It produces cytidine 5'-diphosphocholine (CDPCho) via the S_N2 reaction of choline phosphate (ChoP) and cytidine 5'-triphosphate (CTP)[15,12,16] (Scheme 1). Ligands of *Pf*CCT are bound by several secondary interactions in a highly conserved pocket including a 'composite aromatic box'.[17] The cytosine and pyrophosphate groups are stabilized mainly by hydrogen bonds, while the choline group forms cation- π interactions with Trp692 and Tyr714 residues in the aromatic cage and a salt bridge with the side chain of Asp623 (see Fig. 2). As our primary focus was on cation- π interactions we studied the wild type (WT) enzyme and two experimentally well-characterized mutant enzymes where the original tryptophan residue at position 692 was mutated to tyrosine (Trp692Tyr) and phenylalanine (Trp692Phe), respectively.



Fig. 1. Three-dimensional representation of the homodimeric structure of the *Pf*CCT enzyme. The cytidine 5'-diphosphocholine (CDPCho) ligand is shown in licorice.



Scheme 1. The reaction of choline phosphate (ChoP) and cytidine 5'-triphosphate (CTP) catalyzed by *Pf*CCT producing cytidine 5'-diphosphocholine (CDPCho).



Fig. 2. Most important intermolecular interactions responsible for the binding of cytidine 5'diphosphocholine in the active site of the PfCCT enzyme.

2. Computational details

2.1 Gas phase calculations. Tetramethylammonium ion (TMA) was used as a model of the choline group and its interaction with the side chains of Trp, Phe and Tyr amino acids was investigated. As it is known that most common DFT functionals fail to describe non-covalent interactions properly, we used various functionals parametrized for the better description of non-covalent interactions, the ω B97X-D[18] and the M06-2X[19] functionals. We also calculated the interaction energy with the B3LYP[20] and the TPSSh[21] functionals. For B3LYP the DFT-D3 dispersion correction term[22] was also determined. The structure of the van der Waals complexes of TMA and the amino acid side chains was optimized in vacuo using the ω B97X-D and B3LYP functionals and the 6-31G(d) and cc-pVTZ[23] basis sets using Gaussian09 program.[24] To test the effect of the choice of DFT functional, interaction energies were also calculated using TPSSH and M06-2X functionals at the ω B97X-D/6-31G(d) geometry. The counterpoise correction[25] was used to avoid basis set superposition error. Natural charges were calculated and the NEDA analysis was performed with the NBO 5.9 program.[26,27]

2.2 MD simulation in enzymes. No crystal structure of the *Pf*CCT enzyme exists, therefore a previously published homology model (PMDB: PM0078719)[28] of the native enzyme product(CDPCho) complex was used as a starting structure for our MD simulations. In the case of the point-mutated enzymes, the Trp692 residue was mutated in silico to Tyr and Phe residues. The same protocol was followed in the case of the wild type and mutant enzymes. The protonation state of the ionisable amino acid side chains of the enzymes were predicted by the PROPKA program.[29-32] Based on the estimated pKa values the Glu638 and the Asp589 residues were protonated and His709 was doubly protonated. All other histidine residues were protonated in the ε position based on careful examination of their hydrogen-bond environment. Missing hydrogen atoms were added and minimized according to the standard CHARMM protocol. Buffer region was assigned 21-25 Å away from the alpha-phosphorus atom of the CDPCho ligand, which was set as the centre of the system. Charged residues in the buffer region were neutralized using "patch" residues. The structure was solvated within a 60 Å water box containing TIP3P[33] water molecules and water molecules whose oxygen atom was within 2.8 Å area of other non-hydrogen atoms of the protein or was farther than 25 Å from the alphaphosphorus of the CDPCho ligand were removed. Added water molecules were equilibrated by stochastic boundary MD at 310 K over 20 ps, followed by a multistep optimization of all atoms within a 25 Å sphere around the centre. This was followed by a stochastic boundary MD simulation in which the position of atoms less than 21 Å away from the centre of the system was propagated using Newtonian dynamics, and in the buffer region with Langevin dynamics. First the system was heated to 310 K over 60 ps. At this temperature MD equilibration was carried out over 100 ps, which was followed by a 20 ns productive MD simulation. Increasing restraints were applied in the buffer region in the course of minimizations and MD simulations in order to restrain the movement of heavy atoms around their initial positions. Atoms farther than 25 Å from the alpha-phosphorous atom were fixed. The CHARMM software package[34] and the CHARMM27 force field[35] was used for these simulations and the VMD program[36] for visualizations. Topology file and parameters for the CDPCho ligand were taken from our previous publication.[28] It should be noted that the parameters for the choline group of the ligand are identical to those from the CHARMM lipid force field.[37]

2.3 QM/MM calculations In order to assess the effect of protein flexibility on the results of QM/MM calculations 6 snapshots were taken for the three enzyme variants from the corresponding MD trajectories (from the 0th, 16th, 17th, 18th, 19th and 20th ns) as starting geometries and the geometry of these structures were carefully optimized using molecular mechanics (the CHARMM27 force field) before the QM/MM calculations were started. The CDPCho ligand was represented by a tetramethylammonium (TMA) ion in the quantum mechanically described region (QM region). The QM region also included the closest neighbors of the choline group: the sidechains of the Asp623, Tyr714, Ile740, Tyr741 and of the mutated aromatic residue at position 692 (Trp in the wild type and Tyr and Phe in the mutants, respectively). The covalent bond was formally broken between the C_{α} and C_{β} atoms of each residue and hydrogen-type link atoms were used to satisfy the valences of the carbon atoms in the QM/MM calculations. The charge of the group of the MM atoms directly connected to the QM atoms were set to zero to prevent unrealistic polarization effects. Atoms farther than 20 Å from the alpha-phosphorus atom of the CDPCho ligand were fixed and the charge of atoms farther than 25 Å were set to zero. QM/MM geometry optimizations were carried out at the ωB97X-D/6-31G*/CHARMM27 level using the QoMMMa 8.02 program[38] which couples the output files generated by Gaussian09 and TINKER[39,40] program packages. QM/MM interaction energies were determined at the optimized geometries using the ω B97X-D, B3LYP, TPSSH and M06-2X functionals and the cc-pVTZ basis set.

2.4 Energy decomposition in enzymes. In most force-field based methods the overall interaction energy is calculated as a sum of the pair-interaction energies of the fragments. However, it is also known that in many cases the overall interaction energy cannot be calculated

as a simple sum of the pair-interaction energies. In these cases the non-additive term can be approximated by higher-order energy terms. Especially, the three-body term has been shown to have a significant contribution to the overall interaction energy. E.g. in the case of water clusters the contribution of the three-body term has been shown to account for almost 20% of the overall interaction energy at certain geometries, while at other geometries the three-body term was shown to destabilize the structure and oppose binding.[41] Therefore in the present study we extended a previously proposed energy decomposition method.[42] Accordingly the total energy of the system can be written as a sum of one-, two-, three-, ... n-body terms and the interaction energy (E_{int}) can be calculated according to eq. (1):

$$E_{\text{int}}^{tot} = E_{total} - \sum_{i=1}^{n} E(i) = \sum_{i=1}^{n-1} \sum_{j>i}^{n} \Delta^2 E(ij) + \sum_{i=1}^{n-2} \sum_{j>i}^{n-1} \sum_{k>j}^{n} \Delta^3 E(ijk) + \sum_{i=1}^{n-3} \sum_{j>i}^{n-1} \sum_{k>j}^{n} \Delta^4 E(ijkl)...$$
 (1)

where n is the number of the fragments in the system E(i), E(ij), E(ijk) etc are the energies of the various one, two three membered-clusters in the system, respectively. The pairwise-additive two-body interactions ($\Delta^2 E(ij)$) and the higher three-body ($\Delta^3 E(ijk)$) four-body . $\Delta^4 E(ijkl)$, etc non-additive components are defined as:

$$\Delta^{2}E(ij) = E(ij) - \{E(i) + E(k)\},$$
(2)

$$\Delta^{3}E(ijk) = E(ijk) - \{E(i) + E(j) + E(k)\} - \{\Delta^{2}E(ij) + \Delta^{2}E(ik) + \Delta^{2}E(jk)\},$$
(3)

$$\Delta^{4}E(ijkl) = E(ijkl) - \{E(i) + E(j) + E(k) + E(l)\} - \{\Delta^{2}E(ij) + \Delta^{2}E(ik) + \Delta^{2}E(il) + \Delta^{2}E(ik)\} - \{\Delta^{2}E(ijk) + \Delta^{2}E(ik) + \Delta^{2}E(ik)\} - \{\Delta^{3}E(ijk) + \Delta^{3}E(ijk) + \Delta^{3}E(ikl) + \Delta^{3}E(ijk)\},$$
(4)

This energy decomposition scheme is generally applicable in both quantum mechanics and QM/MM calculations, and could be applied to any systems composed of any fragments. However, in the case of QM/MM calculations some extra considerations are needed. One important note is that in this case it is only worth decomposing the interaction energy of the QM region: in the MM region all energies are calculated from pair potentials. Second consideration regards the type of embedding scheme used for the coupling of the QM and MM regions. In the case of QM/MM calculations with mechanical embedding there is no electrostatic interaction between the QM and MM regions. Therefore the energy decomposition scheme can be applied similarly to QM only calculations and neglecting the term representing the MM energy. However, in the case of calculations with electrostatic embedding care as to be taken with the treatment of the electrostatic interactions. Therefore, we calculated the energy of each fragment (whether one, two, three, four or five or six body), in the presence of the point

charges representing the enzymatic environment but paid attention to eliminate the self energy of the MM point charges from the calculated energy in order to avoid multiple subtraction.

Our QM region comprised of six fragments (the TMA group of the ligand and the five amino acid side chains) and we determined all 3-, 4-, 5- and 6- body energy terms. These calculations were carried out at the ω B97X-D/cc-pVTZ level and included the full set of basis functions used for the description of the entire QM region, in order to avoid the basis set superposition error.

3. Results

3.1 Gas-phase calculations. We started our investigation of cation- π interactions with determining the geometry of model systems of choline and amino acid residues with aromatic side chains. Tetramethylammonium cation (TMA) was used as a model of the quaternary ammonium group of the choline residue and the aromatic amino acids were models with their side chains and their β -carbon atom.



Fig. 3. Optimized structure of investigated models of TMA-Trp/Phe/Tyr complexes at the ω B97X-D/6-31G* level. Selected distances between the quaternary nitrogen atom and the centre of the aromatic rings and the tyrosine oxygen are also shown.

The obtained geometries of the complexes are depicted in Fig. 3. The geometries obtained with the B3LYP and ω B97X-D functionals and the 6-31G* and cc-pVTZ basis sets were very similar. However, the interaction energies between the fragments show very strong functional dependence (see Table 1). Functionals parametrized for improved description of non-covalent interactions (M06-2X and ω B97X-D) predicted significantly stronger interaction between TMA and the aromatic fragments than the TPSSh and B3LYP functional. However, when the D3 dispersion correction is added to the results of the B3LYP calculations, the interaction energy increases significantly, and is in the same range as given by M06-2X and ω B97X-D. It is also worth emphasizing that all functional (whether including or not dispersion correction) predict the same trends for the interaction energies in the studied complexes: phenylalanine forms the weakest cation- π interaction that it is primarily tryptophan that is involved in energetically significant cation- π interactions in proteins.[43]

Table 1. Interaction energies (in kcal/mol) for the gas-phase van der Waals complexes calculated at the ω B97X-D/6-31G* geometries using the cc-pVTZ basis set and the indicated DFT functionals.

			11 551	20211	
TMA-Trp	-17.3	-16.0	-9.1	-7.9	-16.5
TMA-Tyr	-13.8	-12.8	-7.7	-7.0	-13.6
TMA-Phe	-11.7	-10.0	-5.9	-5.23	-11.4

ωB97X-D M06-2X TPSSh B3LYP B3LYP-D3

The NBO analysis[44] has become one of the most significant tools in the hands of computational chemists to gain insight into the nature of interactions in molecules or complexes. We have performed a NEDA analysis[26] of these complexes. In all cases delocalization from the bonding orbitals of the aromatic ring to the antibonding orbitals of C-H bonds located nearest in space to the aromatic ring are observed. Interestingly, in the case of the tyrosine-TMA complex, NEDA indicates that the strongest interaction is observed between the oxygen lone pair and and antibonding orbital of the methyl group of TMA (LP $_{O1} \rightarrow$ BD* c_{3-H32}). This interaction is very well reflected by relative positions of the two fragments: TMA is not positioned strictly over the phenyl ring, but moved closer to the hydroxyl group (see Fig. 3.)

3.2. Molecular dynamics simulations. After having studied the relevant cation- π interactions in gas-phase we turned our attention towards the enzymatic calculations. We did not observe any significant structural distortions of the wild type and mutant enzymes in the course of 20 ns-long MD simulation indicating the good quality of the starting structures. We have monitored the distance between the quaternary nitrogen atom of the choline group and the centre point of the mutated residue (Trp692, Phe692, Tyr692). As Fig. 4A. shows this distance became slightly shorter during the MD in the wild type enzyme, while it remained almost constant in the mutants, showing a stronger interaction in the wild type enzyme. We have also monitored the distance between the nitrogen and the carboxylate oxygens of a nearby aspartate residue (Asp623) (Fig.4B), as the experimental work suggested that it contributes to the binding of the ligand via electrostatic interactions.[14] What is quite apparent is that in the mutants the aspartate-choline distance has decreased which indicates the strengthening of their interaction.



Fig. 4. Distance between the nitrogen atom of CDPCho and the center point of residue 692 (A) and the aspartate oxygens (B) during the MD simulations in the three enzyme variants.

In order to investigate the effect of the protein flexibility on the results of QM/MM calculations, we have taken six snapshots from the trajectories of the three enzyme variants: at 0th ns (thus the last structure obtained at the heating up period), and at the 16th, 17th, 18th, 19th and 20th ns of the trajectories. These structures were chosen because we anticipated that the structures will be fully relaxed after 15 ns of MD simulation and we wanted to compare the results obtained for structures before and after the MD simulation. We carried out a full QM/MM optimization of all snapshots at the ω B97X-D/6-31G*/CHARMM27 level and carried out a full energy decomposition of the interaction energy of the system at the ω B97X-D/cc-pVTZ/CHARMM27 level.

In Fig. 5. we have superimposed the QM/MM optimized geometries of the QM regions of all calculations. It is apparent that compared to the 0th ns structure the structures underwent some changes, and that there is some variability in the structures between 16-20 ns, however most of the changes are most likely due to thermal motion. These figures also show the abovementioned effect: in the wild type enzyme the Asp623 residue went farther away from the ligand then in the mutants.

The experiments on the binding of the CDPCho ligand to the active site of the wild type and mutant PfCCT enzymes showed that the binding in all cases is enthalpy driven (see Table 2),[16] thus it is a reasonable approximation to compare the calculated interaction energies of the ligand within the active site with the experimentally observed heat of association. Still, it is important to note that the experimental value will include other effects than simply the interaction between the ligands and the protein side chains, as it will necessarily include the heat chain related to the distortion of the ligand/protein upon binding as well as the desolvation of the ligand and of the cavity. According to the experiment the ligand binds most strongly to the wild type enzyme, while the mutants show considerably weaker binding, with slight preference for the tyrosine mutant. The gas phase data (Table 1) correctly reproduces this trend and on top, the binding energy between the side chain of residue 692 and the ligand at the starting structure (Ons) and at the last point of the MD (20ns) shows exactly the same trend. The average of the binding energies of the five snapshots (16-20ns structures) confirms that the wild type (Trp) enzyme binds the ligand most strongly, and indicates similar interaction strength in the case of the Phe and Tyr mutated enzymes, in accordance with the experiments. Therefore, we may conclude that the data show little variation, which suggests that a limited number of snapshots may already yield reliable data. Previous studies on other thermodynamic properties such as the S-O[45] and Fe-O bond enthalpies[46] showed also only very small variation.

However, when the overall interaction energy of the ligand with the total QM regions is assessed, the agreement with experiment is not retained, especially not for the 16-20 ns structures, here the wild type enzyme seems to bind the ligand the most weakly. For this reason we have calculated and extra snapshot taken at the 8th ns of the molecular dynamics trajectory as at this point, the Asp623 residue is still located closer to the ligand in the wild type structure. As discussed in the following chapter, the Asp residue is responsible for the majority of the binding energy of the ligand in the active site, and based on our results, this term, as it describes a cation-anion interaction, is very sensitive to the exact position of the moieties. However, it is also likely that due to solvation/desolvation effects this term will be much smaller in reality than predicted by our calculations.



Fig. 5. Superposition of the QM/MM optimized structures of the QM regions in the (A) wild type (B) Trp692Tyr mutant (C) Trp692Phe mutant enzymes. Hydrogen atoms are omitted for clarity. Coloring code for the atoms of the 0 ns structure: light blue: carbon, red: oxygen, dark blue: nitrogen

Table 2. Interaction energies (kcal/mol) of the CDPCho ligand with residue 692 in the enzyme ($E_{int}(e)$) and with the whole QM region in parenthesis, in kcal/mol) according to QM/MM calculations, and experimentally measured enthalpy ($\Delta_a H$ in kcal/mol), entropy ($\Delta_a S$ in cal/molK) and Gibbs free energy ($\Delta_a G$ in kcal/mol) of association at 20 °C. For comparison calculated gas phase data for the interaction of TMA with Trp, Tyr and Phe sidechains is also indicated ($E_{int}(g)$, kcal/mol).

	Eint(e)ons	Eint(e)8ns	Eint(e)16-20ns	Eint(e)20ns	E _{int} (g)	$\Delta_{a}H[14]$	$\Delta_a G[14]$	$\Delta_a S[14]$
Wild Type	-12.5 (-90.4)	-17.0 (-81.4)	$-11.4 \pm 0.8 (-71.4 \pm 3.2)$	-10.8 (-71.5)	-17.3	-22.9 ± 1.0	-5.8 ± 2.5	-58.1 ± 5.0
Trp692Tyr	-8.8 (-93.0)	not calculated	$-6.8 \pm 2.1 \ (-81.5 \pm 3.4)$	-8.7 (-82.8)	-13.8	-9.8 ± 4.5	-4.9 ± 7.0	-16.8 ± 8.6
Trp692Phe	-6.5 (-86.8)	-7.4 (-78.1)	$-7.8 \pm 1.0 (-75.9 \pm 0.5)$	-6.7 (-75.9)	-11.7	-6.8 ± 0.8	-4.3 ± 1.2	-8.9 ± 1.3

3.4. Energy decomposition in enzymes

We carried out a full energy decomposition of the interaction energies $\omega B97X$ -D/cc-pVTZ/MM level. In Table 3 the decomposition of the overall interaction energy to 2-, 3-, 4- and 5-body terms are shown (the contribution of the six-body term is negligible.). The calculated overall interaction energy varies considerably between 80 to 92 kcal/mol, but the percentages of the different terms remain almost constant over all snapshots. Interestingly the two-body term overestimate the binding energy by about 7 kcal/mol, 10% of the overall interaction energy, which is counteracted by the three-body term, which seems to destabilize the interaction. The sum of the four-body terms accounts for 1.7% of the overall interaction energy, but taken the large number of four body terms $\binom{6}{4} = 15$, the contribution of the individual terms is negligible, and those of the five body terms are even smaller. This decomposition shows that the threebody terms contribute indeed significantly to the interaction energy and we observe an overall anticooperative effect as shown by the opposite sign compared to the two-body terms. Interestingly, recently published work on the interplay between various non-covalent interaction found that cation- π interactions acted frequently in an anticooperative manner with other non-covalent interactions e.g. with hydrogen bonds.[47] In Table 4 we have only summarized those terms that are related to the binding of the ligand in the active site in contrast to Table 3 where all terms were summarized (i.e. the interaction energy between two amino acid side chains as well). It is remarkable that 90% of the overall interaction energy is due to ligand binding, and that the ratio of the three body terms is the same as in the case of the overall interaction energy. The same analyses were carried out for the two mutant structures as well, and the same conclusions were drawn (data not shown).

In Table 5 we show the calculated two-body interaction energy terms and those three-body terms that exceed 1 kcal/mol. It is visible that the standard deviation of the most significant terms is much smaller than the value, but in the case of weak interactions quite large standard deviations are observed. It is also obvious that in all cases the anion-cation interaction between the ligand and Asp623 is responsible for the majority of the interaction energy and that this energy is 6-10 times larger than the interaction between the ligand and the aromatic residue at 692. Although, this observation suggests that the interaction with the negative residue is primarily responsible for the strong binding of the ligand in the composite- aromatic box, and these numbers should be considered with some caution, as the desolvation effects upon binding of the ligand in the active site are expected to be more significant in the case of anionic residues

than for aromatic side chains. It is also clear that the closer the aspartate to the CDPCho ligand is the stronger the interaction is (WT: distance 5.4Å E_{int}: -61.8 kcal/mol; Trp692Tyr distance: 4.7 Å E_{int}: -71.4 kcal/mol; Trp692Phe 3.9 Å E_{int}: -85.2 kcal/mol). Interestingly in the case of the Trp692Tyr system the second strongest interaction is found between the Tyr741 and the Asp623 residues. Careful investigation of the structure revealed that a strong hydrogen bond has been established between Asp623 and Tyr741 in this mutant (average distance 1.62 Å). Furthermore a very strong interaction network was established between Asp623, Tyr741 and the ligand as shown by the surprisingly large three-body term (7.3 kcal/mol). Furthermore the data shows that only those three-body terms became stronger than 1 kcal/mol which involved the two charged residues (TMA and Asp623). It is also visible that in all cases CDPCho ligand interacts stronglier with residue 692 than with Tyr741.

Table 3. Decomposition of the total interaction energy (E_{int}^{tot}) and percentage contribution of the various n-body terms to the overall interaction energy for the various QM/MM optimized structures for the wild type enzyme.

time (ns)		16		17		18		19		20	ave	erag	e
Etot	-85.8		-91.9		-89.2		-80.1		-87.2		-86.8	±	4.4
$\Sigma \Delta^2$	-92.3	(107.6 %)	-99	(107.8 %)	-96.6	(108.3 %)	-85.5	(106.8 %)	-92.9	(106.6 %)	-93.3	±	5.1
$\Sigma \Delta^3$	7.7	(9.0 %)	8.4	(9.1 %)	8.5	(9.5 %)	6.3	(7.9%)	6.6	(7.6%)	7.5	±	1.0
$\Sigma \Delta^4$	-1.3	(1.5 %)	-1.4	(1.5 %)	-1.3	(1.5 %)	-1	(1.2 %)	-0.9	(1.0%)	-1.2	±	0.2
$\Sigma \Delta^5$	0.1	(0.1 %)	0.1	(0.1 %)	0.1	(0.1 %)	0.1	(0.1 %)	0.1	(0.1 %)	0.1	±	0.0
$\Sigma\Sigma\Delta^{1-5}$	-85.8	(100.0 %)	-91.9	(100.1 %)	-89.2	(100.0 %)	-80.1	(100.0 %)	-87.2	(100.0 %)	-86.8	±	4.4

Table 4. Decomposition of the interaction energy between the CDPCho ligand and the other fragments ($E_{int,CDPCho}^{tot}$) and percentage contribution of the various n-body terms (compared to the overall interaction energy E_{tot}) for the various QM/MM optimized structures for the wild type enzyme.

time (ns)		16		17		18		19		20	ave	erag	e
$\Sigma \Delta^2$	-77.3	(90.1 %)	-81	(88.2 %)	-81.4	(91.2 %)	-71.7	(89.6 %)	-77.1	(88.4 %)	-77.7	±	3.9
$\Sigma \Delta^3$	7.6	(8.9 %)	8	(8.7 %)	8.4	(9.4 %)	6.3	(7.9%)	6.4	(7.3 %)	7.4	±	0.9
$\Sigma \Delta^4$	-1.2	(1.4 %)	-1.2	(1.3 %)	-1.3	(1.5 %)	-1	(1.2 %)	-0.8	(0.9%)	-1.1	±	0.2
$\Sigma \Delta^5$	0.1	(0.1 %)	0.1	(0.1 %)	0.1	(0.1 %)	0.1	(0.1 %)	0.1	(0.1 %)	0.1	±	0.0
$\Sigma\Sigma\Delta^{1-5}$	-70.8	(82.6 %)	-74.1	(80.7 %)	-74.1	(83.1 %)	-66.3	(82.8 %)	-71.5	(82.0 %)	-71.4	±	3.2

Table 5. Two-body interaction energy terms (in kcal/mol) and three-body terms exceeding 1 kcal/mol with their standard deviations averaged over the 16-20 ns structures for the enzyme variants. Interacting fragments: 1 = Asp623, 2 = Residue692 (Trp/Tyr/Phe), 3 = Tyr714, 4 = Ile740, 5 = Tyr741, 6 = TMA part of the CDPCho ligand.

Interacting Fragments	Wild Type			Trp	692	Tyr	Trp6	Trp692Phe				
12	1.09	±	0.18	-0.52	±	0.68	0.36	±	0.31			
13	0.19	±	0.70	-1.27	±	0.73	-0.33	±	0.28			
14	-0.83	±	0.41	-1.63	±	0.74	-1.36	±	0.27			
15	-7.47	±	0.72	-28.05	±	3.04	-10.40	±	0.20			
16	-61.80	±	3.99	-85.15	±	2.08	-71.42	±	1.35			
23	-1.83	±	0.91	-3.43	±	1.79	-1.42	±	0.45			
24	-0.63	±	0.32	-0.42	±	0.14	-0.80	±	0.17			
25	-0.22	±	0.02	-0.12	±	0.02	-0.09	±	0.04			
26	-11.37	±	0.82	-6.80	±	2.12	-7.84	±	1.00			
34	-2.50	±	0.23	-3.07	±	0.77	-2.75	±	0.55			
35	-0.63	±	0.28	-1.63	±	0.34	-0.53	±	0.23			
36	-3.61	±	1.24	-0.59	±	2.45	-3.84	±	0.94			
45	-2.70	±	0.24	-2.20	±	0.59	-2.64	±	0.15			
46	-0.69	±	0.55	0.05	±	0.44	0.05	±	0.33			
56	-0.25	±	0.69	-1.38	±	1.11	-0.39	±	0.81			
126	2.16	±	0.18	2.61	±	0.38	2.01	±	0.17			
136	1.31	±	0.23	2.02	±	0.20	1.75	±	0.15			
146	1.01	±	0.14	1.28	±	0.37	1.02	±	0.11			
156	2.20	±	0.24	7.32	±	0.72	3.07	±	0.21			

3.5. Assessment of various factors offering possibilities for reduction of computational cost

So far we have established that our QM/MM calculations correctly predict the effect of the studied point mutations on the ligand binding energy of CDPCho and also found that pair potentials account for 90% of the interaction energy. It also became evident that there is very little difference between the studied snapshots taken at 1 ns intervals after proper equilibration of the systems. Therefore we set out to study various other effects on the accuracy of the calculation using the snapshot taken at 20 ns of the MD simulation of wild type.

The calculations presented so far were carried out with a large basis set (cc-pVTZ), so we calculated the pair-wise interaction energies with the smaller 6-31G* basis set, and there was

practically no effect of changing the basis set (see Table 6). This implies that in these QM/MM calculations no large basis sets are necessary; the transferability of this finding to other systems should be tested. Then we turned out attention towards the effect of density functionals. Most of the pair potentials show very weak functional dependence, with the exception of the cation- π interaction between the ligand and the aromatic side chains, although this is most apparent for tryptophan residue 692, as it interacts most strongly with the ligand. Here, B3LYP and TPSSH gives about 9 kcal/mol weaker interaction energy than M06-2X and ω B97X-D, and this value is in perfect accordance with the results of the gas-phase calculations, showing that from simple gas-phase calculations exact functional dependence of the QM/MM results can be deduced.

Table 6. Effect of various factors (QM/MM geometry optimization, basis set, DFT functional, electrostatic embedding of point charges in the QM region, only MM energy calculation) on the obtained two-body interaction energies between TMA and the residues included in the QM region for the wild type structure taken at 20 ns from the MD trajectory. Data is given in kcal/mol.

Wild Type 20ns	asp623	trp692	tyr714	ile740	tyr741
wB97xD/cc-pVTZ/MM	-59.0	-10.8	-5.3	-1.6	-0.5
ωB97xD/6-31G*/MM	-59.3	-11.2	-5.5	-1.7	-0.7
B3LYP/cc-pVTZ/MM	-58.3	-1.9	-3.0	1.5	0.6
TPSSh/cc-pVTZ/MM	-58.5	-3.1	-3.1	1.2	0.6
M06-2X/cc-pVTZ/MM	-58.8	-9.7	-4.7	-0.8	0.0
ωB97xD/cc-pVTZ/MM [§]	-58.2 [§]	-8.3 [§]	-5.1 [§]	-0.8 [§]	-0.2 [§]
E_{strain}^{\dagger}	0.8	1.0	3.0	0.5	0.5
$\omega B97 x D/cc$ -pVTZ [£]	-61.0	-14.1	-7.1	-4.0	-2.5
MM [§]	-17.2 [§]	-15.9 [§]	-6.8 ^{§*}	-0.8 [§]	1.4 [§]

[§] calculated at the MM optimized geometry of the snapshot taken from the MD trajectory ^fgas phase calculation on the QM region

* discrepancy between QM/MM and MM values was much larger for all other structures

^{\dagger} E_{srain} was calculated as the energy difference between the geometry of the side chain adopted in the QM/MM optimized structure and the minimum energy structure in the gas phase One of the most time consuming part of our calculations was the process of QM/MM geometry optimizations. Therefore, we investigated the effect of QM/MM geometry optimization on the interaction energies. We compared the interaction energy calculated at the geometry of the snapshot taken from the MD trajectory and after QM/MM optimization (Table 6 and Table S2 in the Supporting Information). The observed differences were very small, the interaction energies are slightly more favorable at the QM/MM optimized geometry. However this change is so little, that would justify the usage of carefully MM optimized geometries sampled from MD trajectories for QM/MM interaction energy calculations. The very small interaction energy difference calculated for structures before and after QM/MM optimization points to small differences between these structures. Indeed, when we compared the strain energy for each of side chains included in the QM region we observed very small effects (Table 6). The effect is largest for theTyr714 residue reaches 3 kcal/mol.

QM/MM calculations were developed with the desire and need to be able to take into account the effect of the protein environment in course of calculations. However with the development of computing resources one is able to treat larger and larger QM region and now there are advocates of both cluster-based QM calculations on enzymes and QM/MM methods as well, both of which are finding their entries into the pharmaceutical industry as well.[48] Therefore e.g. in the course of modeling enzymatic reactions, if possible, it is worth increasing the size of the QM region to be as large as to have a small effect on the energetic of the reaction, thereby ensuring that all significant interactions are treated at a high level. Too large contribution from the MM region brings up the danger that some essential effects are treated with molecular mechanics thus their description might be unreliable. We checked this and we found that removal of the point charges from the Hamiltonian has a very small effect on the obtained interaction energies, which are slightly stronger in the absence of the point charges.

Based on the above results it seems likely that reasonable interaction energies could be obtained for the interaction of ligands with active site residues from carefully MM optimized snapshots taken from a well-equilibrated trajectory, and that these interactions are already well-predicted from a single structure, without the need to study an ensemble of structures. However, these calculations still require some quantum mechanics based calculations. How well are these data reproduced by molecular mechanics? To answer these questions we have collected all pair-wise interaction energies for all QM/MM optimized structures calculated with the CHARMM27 force field (see Table 6 for the 20 ns WT structure and Table S2 in the Supporting Information for the data of all enzyme variants and all snapshots). It is remarkable that although in the salt-

bridge interaction between TMA and Asp623 is predicted to be the strongest interaction by both methods in all structures, their strength differs, considerably, by 20 kcal/mol. The second most important interaction is the cation- π interaction between TMA and residue 692, however the MM method overestimates this interaction by 6-9 kcal/mol, which exceeds by 50% the QM/MM interaction energy. The third most important interaction is found between TMA and Tyr714, and both methods predict it to be around 3-5 kcal/mol, which seems to be consistent with the cation- π interaction energy strength calculated in the gas phase complexes (~14 kcal/mol), as the distance in the enzymatic pocket (~5 Å) is much larger between the fragments than in the gas-phase van der Waals complex (~4 Å). The interaction between IIe740 and Tyr741, and TMA are predicted to be very weak by both methodologies. The discrepancy observed between the QM/MM and MM interaction energies is in line with a recent study on protein-ligand binding energies that found 15% difference between MM and QM energies even after adjustment for systematic differences.[49]

The significant discrepancy between the calculated MM and QM/MM interaction energies suggest that the parameters of the choline group (that were taken from the CHARMM lipid force field[37]) should be treated with great care and may require further validation for enzymatic systems. We also note here, that the original parameters were developed for the aim of reliable modeling of membranes, which may be the underlying reason that in the present simulation so large differences were found.

Conclusions

In the present study we investigated cation- π interactions using a variety of methods focussing on the key enzyme of the membrane synthesis (phosphocholine cytidylyltransferase) of the pathogen of malaria. We have adopted a methodology for the computation of many-body interactions in proteins using QM/MM and showed that in our case the two-body terms overestimate the overall interaction energy of the ligand by about ten percent, while three-body interaction terms, which are dominated by cation- π interactions, counteract the two-body terms and by decreasing the strength of the other interactions. Using accurate QM/MM calculations of the binding energy we obtained good agreement between the trends of the binding affinities of the wild type and mutant enzymes. Furthermore we tested the effect of various factors that could affect the accuracy and reliability of the calculations and sought approximations that could offer reduction in computing time without reducing reliability. We found that once well-equilibrated structures

were reached in the MD simulations, very similar results can be obtained for the various snapshots taken from the trajectory. The calculated interaction energies were only slightly influenced by electrostatic embedding of the point charges in the QM/MM calculations, and by QM/MM geometry optimization. However, the calculated molecular mechanics interaction energies were off by more than 50% for cation- π interactions involving a choline group and even large discrepancies were observed for its interactions with an acidic side chain. Therefore, careful validation of the force fields for the estimation of cation- π interaction energies is highly recommended before using the force field for virtual screening seems. Based on our results we suggest to calibrate force fields based on fragment-based QM-calculations on geometries obtained from MD simulations as this approach is likely to yield reliable binding energies at reduced computational cost.

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