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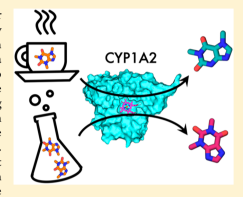
Binding Modes and Metabolism of Caffeine

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Supporting Information

ABSTRACT: A correct estimate of ligand binding modes and a ratio of their occupancies is crucial for calculations of binding free energies. The newly developed method BLUES combines molecular dynamics with nonequilibrium candidate Monte Carlo. Nonequilibrium candidate Monte Carlo generates a plethora of possible binding modes and molecular dynamics enables the system to relax. We used BLUES to investigate binding modes of caffeine in the active site of its metabolizing enzyme Cytochrome P450 1A2 with the aim of elucidating metabolite-formation profiles at different concentrations. Because the activation energies of all sites of metabolism do not show a clear preference for one metabolite over the others, the orientations in the active site must play a key role. In simulations with caffeine located in a spacious pocket above the I-helix, it points N3 and N1 to the heme iron, whereas in simulations where caffeine is in close proximity to the heme N7 and C8 are preferably oriented toward the heme



iron. We propose a mechanism where at low caffeine concentrations caffeine binds to the upper part of the active site, leading to formation of the main metabolite paraxanthine. On the other hand, at high concentrations two molecules are located in the active site, forcing one molecule into close proximity to the heme and yielding metabolites theophylline and trimethyluretic acid. Our results offer an explanation of previously published experimental results.

■ INTRODUCTION

Human cytochromes P450 (CYP) are oxidoreductases with a heme cofactor that are responsible for the Phase I metabolism of 75% of drugs in the human body. 1-3 There are 57 mammalian isoforms known and their inhibition, induction, or allosteric effects by various small molecules often lead to a number of drug-drug interactions. Cytochromes P450 catalyze a wide variety of reactions, that are in general improving the water solubility either of their endogenous substrates or xenobiotics. These reactions take place at the functional groups of the molecules, also known as sites of metabolism (SOM). A general rule-of-thumb is that poses for which the distance between a SOM of a molecule to the heme iron is not more than 6 Å are considered to be active binding modes.^{4,5} Here, we chose the 1A2 isoform, which metabolizes caffeine in four positions. Caffeine is a methylxanthine neurostimulant, acting as a competitive antagonist of adenosine receptors, that most Europeans consume every day. 6-8 Caffeine, like many other aromatic and heterocyclic amines, is metabolized by CYP1A2.9 CYP1A2 has a relatively narrow and planar binding site (375 Å³) suitable for accommodation of such amines. The active site is formed by the I-helix situated above a cysteine-bound heme with residues Phe226 and Asp320 playing a crucial role in the kinetics of the chemical reaction. 10,11

The exact binding modes of caffeine are unknown, however the main metabolites are known and NMR studies investigating the binding modes and their ratios have been reported. 12 The main metabolites of caffeine are paraxanthine (PX) accounting for about 80% of product formation, theobromine (TB) with about 10%, theophylline (TP) with about 5%, and 1,3,7-trimethyluretic acid (TMU) with only 1%, all of which are biologically active and further metabolized by cytochrome P450s, 13-216 shown in Figure 1.

Regal and Nelson 12 observed a shift in the metabolite ratio

with increasing caffeine concentration, making theophylline the main metabolite at higher caffeine concentrations. CYP1A2 in its resting state is in the high spin state (HS) which is typically associated with a penta-coordinated heme iron in its ferric form.¹⁷ For other CYPs, the resting state involves coordination of a water molecule as sixth ligand to the heme iron, leading to a (measurable) low spin state (LS). The lack of a sixthcoordinating water molecule in the resting state of CYP1A2 might be caused by the narrow hydrophobic active site of CYP1A2, which might to some degree hamper coordination of a water molecule with the heme iron.

A potential substrate binds to the active site in the resting state, potentially in a fixed orientation. For CYP1A2, it was shown that in the presence of higher concentrations of substrate there was an incomplete shift (\sim 28%) from the high spin to the low spin state. 18 Regal and Nelson 12 showed that average distances of caffeine SOMs to the heme iron are approximately 2 Å shorter for CYP1A2 in a 100% low spin

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Figure 1. Main metabolites of caffeine. Caffeine is in the middle and the SOM are depicted in colorful spheres. The main metabolite PX is shown in cyan.

state than in a 100% high spin state. Nevertheless, the distances of all three nitrogens N1, N3, and N7 to the iron atom differ less than 0.2 Å in both spin states, see Table S1. Similar distances of all SOMs to the heme iron do not indicate a clear binding mode leading to the known main metabolites in human

After substrate binding, diffusion of molecular oxygen to the heme iron leads to the formation of compound one (CPDI) with one oxygen atom bound to the iron in its ferryl state. CPDI is a highly reactive species responsible for product formation.

The aim of our work is to elucidate the binding modes of caffeine with the CYP1A2 isoform by using the package BLUES, which can by NCMC lead to a better sampling of ligand orientations.

Activation Energy. The activation energy of the oxidation reaction at each SOM determines the reactivity. Previous studies show that the rate-limiting step in aliphatic oxidation is the creation of a radical intermediate, that is, hydrogen abstraction. The rate-limiting step for aromatic oxidation is the creation of the bond between the oxygen bound to the iron and the aromatic carbon. Rydberg and co-workers ¹⁹ proposed a method predicting these activation energies for potential SOMs based on the effect of their surrounding atoms. This method consists of a number of rules that are derived from high-level density functional theory (DFT) calculations. The calculated activation energies from Rydberg et al.¹⁹ are shown in Table 1. The aliphatic oxidation on N7–C14 and aromatic oxidation on C8 yielded the lowest activation energies, both 52

Table 1. Activation Energies for SOM of Caffeine Taken from Rydberg et al.¹⁹ at Given SOMs^a

SOM	E [kJ/mol]
Aliphatic Oxidations	
N1-C10 (TB)	61.7
N3-C12 (PX)	57.3
N7-C14 (TP)	52.1
Aromatic Oxidation	
C8 (TMU)	52.2

^aAbbreviations of the formed products in brackets.

kJ/mol. The aliphatic oxidations on N3–C12 and N1–C10, were slightly higher, ranging from 57 to 62 kJ/mol.

Despite the lower activation energies of N7–C14 and C8, the overall energies differ by less than 10 kJ/mol, which does not imply a distinctly higher reactivity of one SOM over the others. Apparently, the activation energies alone do not explain the preferred formation of PX (metabolism at N3–C12) but rather suggest a preference for TP or TMU. Caffeine's orientation with respect to the heme might have a stronger impact on its metabolism than its intrinsic reactivity.

Ligand Sampling. Enzymes catalyze chemical reactions typically through a stabilization of the transition state in the active site. The active site might be located on the surface of the protein but might be also buried in the protein interior. In those cases, a ligand or a substrate is steered by nonbonded interactions through the protein along its binding path, passing by potential subpockets until it reaches a reactive binding mode in the protein active site. Assessing the correct ligand binding modes and the ratios of their occurrences is essential for the estimate of ligand binding strength and, for enzyme—substrate complexes, the proper orientation of the substrate.

Binding modes play an important role in drug design where one aims to predict the binding characteristics of a large number of potential drug molecules to the target of interest. Probably the most frequently used and the fastest method for selection of the most likely binders is molecular docking. Molecular docking positions a ligand into multiple binding modes and selects the one with the lowest score (hopefully roughly corresponding to a binding enthalpy) which can be calculated in various ways.²⁰ Docking can reach high computational speed by neglecting the protein motion and solvation. These factors have, however, a negative impact on the accuracy of the results, which makes docking a "guiding filter" to refine ligand libraries and select potentially active binders. Docking does not accurately predict free-energies and occupancies of different binding modes because of the lack of physically accurate sampling.²¹ There are several more extensive methods, employing molecular dynamics (MD) simulations, that address these topics, such as replica exchange, ^{22–24} umbrella sampling with potential of mean force ^{25–27} and others. ^{28–30} These methods are, however, often rather laborious and need a significant amount of preparation for individual systems. Alchemical methods that focus on the relative free energies between individual ligands are often initiated from a single binding pose for multiple ligands. This way they ignore possible orientational differences between ligands which might lead to higher errors and inconsistencies with simulations that start from a different binding pose.^{31,32}

A recently developed method, binding modes of ligands using enhanced sampling (BLUES),³³ addresses these issues. The uniqueness of the BLUES method lies in the combination of MD with a nonequilibrium candidate Monte Carlo (NCMC) approach. The NCMC approach is based on the naïve Monte Carlo method while yielding higher acceptance probabilities with shorter correlation times.³⁴ NCMC introduces a nonequilibrium phase, which consists of a number of instantaneous perturbation steps separated by short relaxation periods. Afterward the entire series is accepted or rejected. In the particular implementation of NCMC used in BLUES, the perturbations scale down the interactions, that is, the potential energy of the ligand with its surroundings is gradually changed to zero, followed by a ligand rotation around its center of mass (COM) into a new orientation without

changing the potential energy of the system. After the ligand rotation, the interactions are gradually turned on again (Figure 2). Coulombic interactions are turned off before the van der

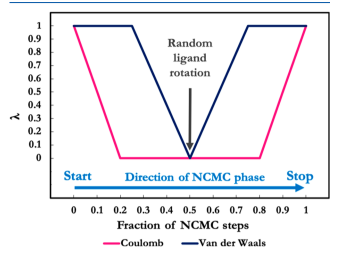


Figure 2. Scaling of nonbonded interactions of the ligand with respect to λ over the course of NCMC steps. At $\lambda=1$ the ligand is fully interacting, whereas at $\lambda=0$ the interactions of the ligand are turned off completely. The relaxation MD part is not included in this figure. Adapted from Gill et al.³³

Waals interactions, and after rotation interactions are turned on in reverse order to avoid numerical instabilities and to ensure reversibility.³⁵

The acceptance criterium for the entire NCMC procedure is derived from the Metropolis-Hastings criterium.³⁶ The acceptance probability of process X, A[X] is computed as

$$A[X] = \min\{1, e^{-w(X)}\}\tag{1}$$

where w is the protocol work of the procedure, estimated as

$$w(X) = \sum_{t=1}^{T} \left[u_t(x_t) - u_{t-1}(x_t) \right]$$
 (2)

where x_t is a microstate at a given simulation step and u_t is the reduced potential energy. Because a Langevin integrator is used, w in eq 1 is complemented with the appropriate shadow work.³⁷ After each accepted or rejected move, the momenta are randomly reassigned based on the Maxwell–Boltzmann distribution in order to keep detailed balance.³⁸ The NCMC stage is followed by a series of conventional MD steps, using a Langevin integrator to relax the entire system. BLUES creates a trajectory of MD and accepted moves, which can be subsequently clustered and used for estimating populations of binding modes.

■ METHODS

The crystal structure of CYP1A2 with the PDB code 39 with a resolution of 1.95 Å was used as an input structure for all MD simulations. The crystallized ligand α -naphthoflavone was subsequently removed and caffeine was docked into the active site using the AutoDock Vina tool and AutoDock Vina scoring function. The active site was defined as a cube with an edge length of 13 Å, centered at the heme iron. Figure S1 shows the docking setup for both receptors. The flexibility of the receptor was not considered. The nine best scored docking poses were saved for further simulations.

Hydrogens were added to the protein using tleap from AmberTools17^{41,42} and the protein was parametrized using the

ff99SB force field.⁴³ The parameters for heme in the ferric form, compound I, and neighboring cysteine were adopted from Shahrokh et al. 44 In a classical force field, no differences between ferric high spin or low spin states are described. Hydrogens were added to caffeine using AutoDockTools445 and caffeine was parametrized using the GAFF⁴⁶ force field with AM1-BCC charges.⁴⁷ The docked proteins were solvated in a rectangular box with TIP3P water molecules 48 with 13 Å as the minimal protein-wall distance and 6 Cl⁻ counterions were added. The system was minimized and equilibrated for 6.2 ns using OpenMM 7.1.1⁴⁹ under constant pressure at 1 atm with 300 K. A Monte Carlo barostat and Langevin integrator were used with a 4 fs time step while setting the hydrogen mass to 3.024 Da and using 1 ps⁻¹ collision rate. The bond distance between hydrogen and heavy atoms was constrained while using the hydrogen mass repartitioning scheme. 50 The particle Mesh Ewald method 51 was used to calculate the long-range electrostatics.

To generate reasonable starting structures for the simulations with the ferric heme, all poses from the last 6 ns of equilibration were merged and analyzed by time-structure-independent component analysis (tICA) using PyEMMA 2.5.2.^{52,53} tICA was performed on the pairwise-distances of the caffeine atoms and the alpha carbons of the binding site with a lagtime of 200 ps and the frames were clustered by PCCA. 34,55 Four of the five most stable clusters were used for standard MD and BLUES simulations. Note that the sole purpose of the tICA analysis and clustering was to obtain initial structures for the production simulations; from such short simulations, no conclusions with respect to the orientational preferences of the substrate can be deduced. Although tICA is also used in the context of building Markov State Models for analysis of long MD simulations, that was not our focus here; instead, we simply sought to identify potential stable or metastable binding modes for further analysis via additional simulations. For the CPDI simulations we did not perform clustering but directly used the rather diverse original docking poses for standard MD and BLUES simulations.

The production runs of standard MD with ferric heme and CPDI were performed by OpenMM in the NVT ensemble without use of a barostat. We have found that running BLUES with a barostat can impair acceptance rates and, for a buried binding site like this, ligand placement is unlikely to affect the pressure of the system. For consistency, the standard MD simulations were also performed without a barostat. In all cases, equilibration brought the systems to the correct pressure prior to these production runs. The production simulations were performed for 1 μ s per docked pose.

BLUES version 0.1.3 was used. The BLUES calculations were performed using openmmtools $0.13.0^{56}$ to annihilate caffeine Coulombic and van der Waals interactions. The MD part was performed as described above by OpenMM's Langevin integrator and the NCMC part by the BAOAB integrator for Langevin dynamics.⁵⁷ The protein and water residues further than 5 Å from the ligand were constrained during the NCMC phase (though these are fully flexible during the conventional MD phase) to improve acceptance, as in previous work.³³ For every perturbation step performed, three additional propagation (or relaxation) steps were applied between λ = 0.2 and 0.8. BLUES runs consisted of 1000 iterations, each of which consisted of 10 000 NCMC steps and 10 000 MD steps. Thus, the total simulation time of one BLUES run is equivalent to 20×10^6 force evaluations. The general scheme of the BLUES workflow is depicted in Figure S2.

■ RESULTS AND DISCUSSION

In order to investigate binding modes of caffeine we included two heme states of the CYP1A2 catalytic cycle. The first of them was heme in its ferric resting state to which the substrate binds and the second was CPDI, which plays a key role in the actual substrate metabolism.

Heme with Ferric Iron. After the removal of the α -naphthoflavone molecule from the crystal structure, caffeine was docked into the active site of CYP1A2. The docking

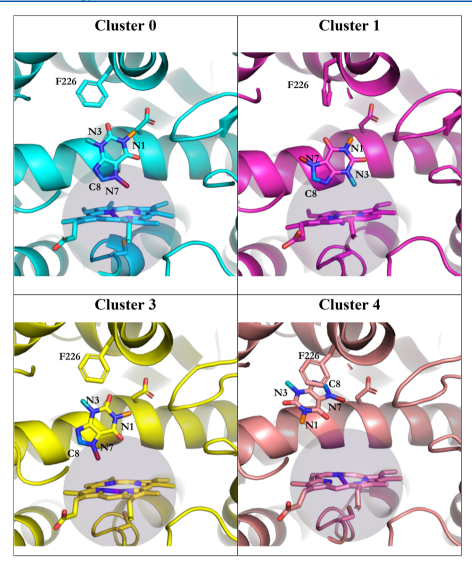


Figure 3. Starting poses for MD and BLUES simulations. The amino acids in sticks are he226 and Asp320, the purple sphere represents a sphere of 6 Å around the heme iron. Caffeine SOMs C10 is in orange, C12 is in cyan, C14 is in dark red, and C8 is in marine blue. The nitrogen atoms to which C10, C12, and C14 are bound, as well as C8, are labeled as well as Phe226.

process resulted in multiple docking poses where at least one functional moiety of caffeine would fulfill the 6 Å rule. Nine well-scoring poses were selected for further simulations. After the minimization and equilibration of docked poses, the feature coordinates were transformed using tICA and subsequently assigned to clusters using PCCA. From five suggested clusters, four were chosen, because the representative structures of clusters 2 and 3 were, based on visual inspection, almost identical. The remaining four clusters were used to start standard MD and BLUES simulations and are shown in Figure 3.

We measured distances of the four sites of metabolism, depicted as C10 (neighboring N1), C12 (neighboring N3), C14 (neighboring N7), and C8, as well as the running average of the distance of the center of mass of caffeine depicted as COM, all to the heme iron of the ferric state.

For the sake of clarity we compare in Figure 4 these distances during standard MD and BLUES simulations of cluster 1 and 4, as the behavior of caffeine in simulations starting from clusters 0 and 3 resemble the data in cluster 1 (Figure S3). For clusters 0, 1, and 3, in BLUES we observed

binding modes consistent with a clear preference for metabolism at C8 and C14, because these SOMs were pointing to the heme iron for more than 90% of simulation frames, see Table S3 and Figure 5. This trend can be seen in MD simulations as well; however, in clusters 1 and 3 caffeine tends to leave the close vicinity of the heme and reaches distances around 10 Å from the heme iron. Because in BLUES the trajectory is divided by the NCMC phase, caffeine does not have the chance for bigger translational moves and remains at a constant distance around 7 Å from the heme. Table S2 shows the percentages of frames, in which the distance between the caffeine SOMs and the heme iron is less than 6 Å. In BLUES simulations, we could not observe any transitions between different binding poses, leading to an average acceptance rate of ~3% in all cluster simulations. Table S1 summarizes average distances of the carbon atoms from the heme iron as $\langle r \rangle$ and as $(r^{-6})^{-1/6}$ which are more representative to the distances that are derived from NMR experiments.

Unlike simulations of clusters 0, 1, and 3, simulations of cluster 4 started from a position where the COM of caffeine was 10 Å from the heme iron. In BLUES simulations of cluster

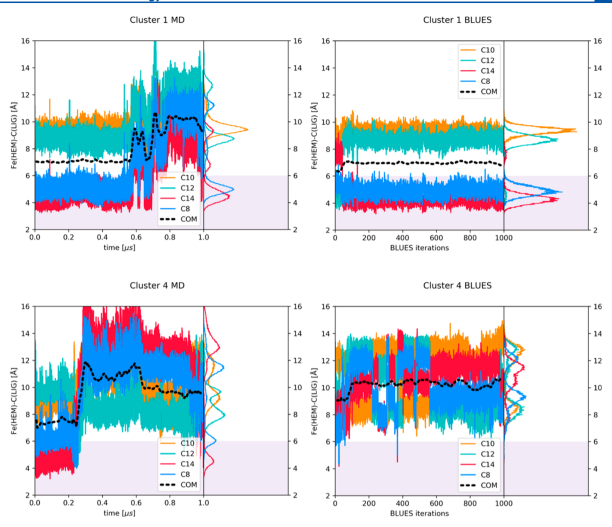


Figure 4. Time series of the distance from the heme iron (ferric state) to indicated atoms in caffeine and its center of mass (COM). The right-hand-side panels show the corresponding histograms.

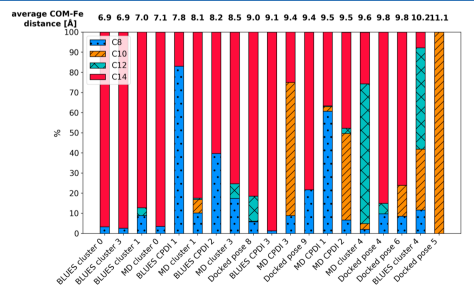


Figure 5. Ratio of the closest SOMs to the heme iron in all simulations. The bars are ordered according to the average COM-Fe distance. We can observe an orientational change in the position where the COM-Fe distance is larger than 9 Å. The exact values and error estimates are shown in Table S7.

4, we observe a higher number of transitions, repeatedly placing all four SOMs of caffeine toward the heme, shown in

Figure 4. In MD simulations of cluster 4, caffeine first comes closer to the heme to around 8 Å, then leaves the active site

and rotates. In both simulations, MD and BLUES, caffeine reaches a metastable pose where C12, which is the methyl group that when cleaved yields the main metabolite PX, is oriented toward the heme iron in more than 50% of simulation frames. It is important to note that in both simulations of cluster 4, C12 does not reach the distance of 6 Å to the heme iron, which is considered to be necessary for the SOM metabolism to take place.

To obtain a better view of the different binding modes, we measured distances from the center of mass of the aromatic cluster (Phe226, Phe256, and Phe260) and the ligand, see Figure S6. We note that when caffeine leaves the close vicinity of the heme, which is the case for clusters 1, 3, and 4, it approaches this aromatic cluster. Favorable aromatic interactions can be the cause of caffeine movement during the MD simulations.

Several factors may cause the differences between BLUES and MD simulations. In BLUES simulations, caffeine does not leave the close vicinity of the heme, because the simulation is constantly interrupted by a series of NCMC steps where it can adjust its orientation. The MD simulations on the other hand, offer a long continuous run where the caffeine molecule has a chance to diffuse away from the direct vicinity of the heme, from which we learn that a single caffeine molecule prefers to bind further away from the heme. The strength of BLUES, however, lies in the effective sampling of all relevant binding modes, where it surpasses standard MD simulations (e.g., cluster 4 in Figure 4).

Measured distances of MD simulations of four additional docked poses that were not used in BLUES are in the Supporting Information (Figure S4), most of them with C14 and C8 oriented toward the heme.

Compound I. Starting poses for simulations with Compound I (CPDI) were obtained from docking using the same procedure as described for the ferric state of the enzyme. These were, however, not clustered, yet the three most distinct docking poses, each orienting a different SOM (C10, C12, and C14) to the heme iron, were chosen for further study. While the binding of the substrate takes place with the enzyme in the ferric resting state, the actual reaction is performed by CPDI. Therefore, we investigated if CPDI affects the orientation of caffeine. During the equilibration run all of them reoriented such that C8 and C14 were pointing to the heme iron. In these MD simulations, the COM of caffeine was at a distance of around 9 Å from the heme iron and C8 and C14 were at 7-8 Å, which was again higher than in BLUES, where these average distances were 8 and 5-6 Å, respectively, see Figure S5 and Table S4. In contrast to the simulation with the ferric iron, the standard MD CPDI simulations showed distances of the SOMs to the heme iron smaller than 6 Å for less than 9% of the frames (Table S5). Interestingly, we observe a stronger preference for C8 than in the ferric heme simulations for both standard MD and BLUES (see Table S6). However, these SOMs are located closely to each other suggesting only a minor difference between heme and CPDI poses. The larger distances are probably caused by the oxygen atom bound to the heme iron and a reduced net charge of the iron-center. These larger distances from the heme iron are correlated with shorter distances to the aromatic cluster (Figure S6).

Figure 5 summarizes the percentages of simulation frames in which each SOM was closest to the heme iron for all simulations. The simulations are ordered by the distance of the COM of the ligand from the heme iron. In binding poses

where the COM is located more than 9 Å from the heme iron, a clear shift toward C12(N3) and C10(N1) is observed. This might suggest that for the binding poses, where the ligand is located further from the heme iron, the metabolite distribution changes and favors formation of PX and TB, while at closer distances TP and TMU are expected as main metabolites.

Two Spin States—Two Mechanisms? The work of Regal et al. 12 suggests a more complicated picture of caffeine dynamics onto which our simulations may shed light. At low caffeine concentrations, PX is the main metabolite and the protein is observed to be in a high spin state. They determined an average distance for all aromatic nitrogens at roughly 7.3 Å, suggesting a favored binding mode that is too far from the heme iron to explain metabolism. Indeed, in many of the MD simulations that start with the caffeine close to the heme, it is seen to move further away, toward the aromatic cluster (Figure 4, Figure S6, Tables S1-3). In the MD and BLUES simulations that place the caffeine at a larger distance, we observe orientations of the substrate, which point C12 toward the heme iron for appreciable amounts of time which is in agreement with the experimentally observed product formation. This binding mode does not only provide caffeine favorable interactions with Phe226 via π – π stacking, see Figure S6, but also places it in a more spacious cavity above the I-helix giving it a higher conformational freedom to rotate.

At high caffeine concentrations, Regal et al. 12 observe a shift in the product formation and in the spin-state of the enzyme to 72% high spin and 28% low spin, see Table 2. We propose that

Table 2. Observed Product Formation (Normalized) As Described by Regal et al.¹² at 5 and 25 mM Caffeine^a

normalized product formation				
SOM	5 mM (100% HS)	25 mM (72% HS + 28% LS)	100% high spin	100% low spin
N1-C10 (TB)	0.11	0.22	0.11	0.25
N3-C12 (TX)	0.78	0.17	0.78	0
N7-C14 (TP)	0.12	0.56	0.12	0.69
C8 (TMU)	0	0.05	0	0.07

^aWe propose that the data at 25 mM with an increased overall turnover by a factor 2.1 is the weighted average of an HS product formation and an LS product formation at fivefold increased activity.

the 28% of low spin corresponds to a situation in which two caffeine molecules bind to the active site forcing one molecule closer to the heme. In Table 2, we compute the normalized product formation for the 100% low spin state as the distribution that would lead to the observed product formation at 25 mM, taking into account the mixed spin state and increased activity (approximately by a factor of 2.1¹²) at the higher concentration. For a 100% low spin state, a 69% preference for product TP with C14 pointing toward the heme and a 5-fold increase in activity relative to the 100% high spin state would be necessary to lead to the observed product formation.

In Figure 6, we summarize the renormalized metabolite yields corresponding to different spin states in Table 2 and compare them to the percentages of the closest SOM to the heme iron throughout our simulations. The main SOM of caffeine in CYP1A2 in its high spin state is N3, which is also the SOM oriented toward the heme iron for more than 50% of the time in both MD and BLUES simulations. In the low spin state, which is present during 28% of time at high caffeine

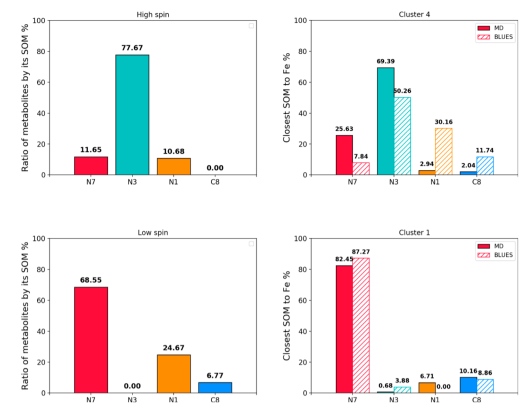


Figure 6. (Left) Renormalized metabolite formation at different spin states, where the bars indicate which SOM has to be the closest to the heme in order for the SOM's metabolism to take place. (Right) Percentages of the corresponding orientation of caffeine during MD (full) and BLUES (striped) simulations. The labels above bars show the percentages of corresponding orientation rounded to two decimal places.

concentrations, the main SOM is N7, corresponding to the most preferred SOM in the simulations, where caffeine is in close vicinity to the heme.

Summarizing, a single substrate molecule in the active site prefers to bind close to the aromatic cluster and relatively far away from the heme group. The orientations that are observed for simulations at this distance agree with the experimentally observed product formation at low concentrations. At higher concentrations, a spin shift is experimentally observed, resulting from a shorter substrate-heme distance. Simultaneously, a shift in the product formation is observed. The orientation of the substrate in simulations at shorter distances agrees with this altered product formation. We propose that a second caffeine molecule in the active site could be the cause for placing the first substrate molecule closer to the heme iron.

MD Simulation with Two Ligands in the Active Site. In order to explore the possibility of simultaneous binding of two ligands in the active site of CYP1A2, we simulated such a complex with standard MD for 1 μ s. The initial structure was created from the initial binding poses of caffeine in cluster 1 and cluster 4 with the enzyme in the ferric state. Both structures were overlaid and the caffeine molecule in the binding mode from cluster 1 was added to the starting structure of cluster 4. This approach was preferred over a new ab initio docking of the two molecules, as, for example, described for Aflatoxin B1 in Cytochrome P450 3A4, sa the two poses represented by clusters 1 and 4 were naturally observed in MD simulations previously and seemed to be readily accommodated in the structure. The system was initially minimized in MOE, sa and subsequently the complex

was minimized, equilibrated, and simulated as described above for the simulations of CYP1A2 with one ligand.

In the production run, both ligands were very stable; the COM of the caffeine molecule closer to the heme iron (Caffeine 1) was located 7 Å from the heme iron whereas the COM of the second molecule (Caffeine 2) was located around 13-14 Å from the heme iron. These distances did not change significantly during the simulation, see Figure S7. Caffeine 1 in the beginning of the production run rotates which results in an orientation that resembles simulations of clusters 0, 1, and 3, that is, with C8 and C14(N7) pointing toward the heme iron. Caffeine 1 stays in this orientation for more than 98% of simulation time with the C8, C14(N7) within 5 Å from the heme iron (Tables S2-3). Caffeine 2 is stabilized by aromatic stacking with Phe226 and Phe260 from both sides and Phe256 from the top. It remains in the same orientation throughout the simulation. The importance of aromatic interactions in CYPs is demonstrated in the defense mechanism of Antarctic sponges.⁶⁰ Here, two small molecules inhibit CYP315a1 and CYP314a1 by aromatic interactions within the active site. Figure 7 shows a snapshot from this simultaneous binding of two caffeine molecules into CYP1A2.

The stability of the protein conformation with two ligands bound was monitored in order to investigate possible effects on the protein upon binding of two caffeine molecules. Figure S8 shows the backbone root-mean-square-deviation (RMSD) curves for this simulation and for the simulations of cluster 1 and cluster 4. Despite the steeper increase of the RMSD curve of protein with two ligands bound, the overall value does not exceed the RMSD from the simulations starting from Cluster 1 and 4. This suggests that simultaneous binding of two caffeine

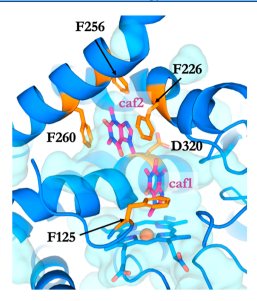


Figure 7. A snapshot of the simulation of simultaneous binding of two caffeine molecules to CYP1A2 after 1000 ns. Caffeine molecules are in magenta, and residues accommodating the caffeine molecules are in orange.

molecules does not have a negative impact on the CYP1A2 stability and in fact may be thermodynamically reasonable. A similar simultaneous binding of two pyrene molecules to the active site of CYP1A2 has been previously suggested by Sohl et al. 61,62

Under physiological conditions in humans, much lower concentrations of caffeine ($<100~\mu\mathrm{M}$) are usually present, and paraxanthine remains the main metabolite, supporting the proposed mechanism with one caffeine molecule in the active site. On the other hand, a double occupancy of the active site might be responsible for the caffeine-induced enhanced metabolism of several druglike compounds. Cameron et al. described heterotropic cooperativity of acetaminophen and caffeine binding in CYP3A4, leading to enhanced metabolism of caffeine. Multiple molecules binding to the active site of CYP3A4 has been described for ketoconazole experimentally and computationally and was postulated for aflatoxin B1 based on observed homotropic cooperativity.

CONCLUSION

In this work, we investigated the binding modes of caffeine to its metabolizing enzyme CYP1A2 via standard molecular dynamics and the enhanced ligand sampling method BLUES. We observed a clear preference of orientation of the N7–C14 and C8 sites of metabolism toward the heme iron, when the center of mass of the ligand was in the vicinity of the active site. This is in agreement with previously published experimental results showing an altered rate of metabolism and shorter caffeine—iron distances at high caffeine concentrations. Preference of N7–C14 and C8 metabolism is also expected from quantum mechanically derived activation energies from literature, which were slightly lower than the activation energies for other SOMs.

In case of caffeine bound further from the heme, we could see a more pronounced conformational freedom due to a larger cavity, where BLUES was able to sample binding poses with all SOMs oriented toward the heme. Additionally, this was the only case where we could observe the N3–C12 moiety facing

the heme iron for a significant amount of time. This binding mode still requires the substrate to transiently approach the heme before metabolism takes place. We suggest that the further removed binding mode corresponds to the binding mode of caffeine when it is present in lower concentrations, where the main metabolite, paraxanthine, is formed. We propose a mechanism, where at higher concentrations two caffeine molecules are accommodated in the active site, forcing one caffeine molecule to be in close vicinity (<6 Å) of the heme iron, leading to formation of theophylline as the main metabolite. To confirm our theory, we simulated simultaneous binding of two caffeine molecules bound to CYP1A2, which indeed lead to stable binding poses with N7-C14 and C8 SOM oriented toward the heme. Moreover, such binding did not disrupt the overall stability of the protein and both caffeine molecules were very stably interacting with the key active site residues. Taken together, the work of Regal et al.,12 complemented by the simulations described in this work, draw a complex but consistent picture of caffeine metabolism by CYP1A2, which might shed more light on caffeine-drug interactions in humans.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.chemrestox.9b00030.

Docking setup for ferric heme and Compound I; BLUES scheme; time series of heme iron to SOM distances for additional simulations; time series of distances from the substrate to the aromatic cluster; atom-positional root-mean-square deviations for selected simulations; average distances and preferred orientations in the various simulations; orientational preferences for all simulations (PDF)

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Notes

The authors declare no competing financial interest.

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