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Insights into the Mechanism of Oxidation of Dihydroorotate to Orotate Catalysed by Human Class 2 Dihydroorotate Dehydrogenase: A QM/MM Free Energy Study

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The Dihydroorotate Dehydrogenase (DHOD) enzyme catalyzes the unique redox reaction in the de novo pyrimidines biosynthesis pathway. In this reaction, the oxidation of dihydroorotate (DHO) to orotate (OA) and reduction of the flavin mononucleotide (FMN) cofactor is catalysed by DHOD. The class 2 DHOD, to which the human enzyme belongs, was experimentally shown to follow a stepwise mechanism but the data did not allow the determination of the order of bond-breaking in a stepwise oxidation of DHO. The goal of this study is to understand the reaction mechanism at molecular level of the class 2 DHOD, which may aid the design of inhibitors that selectively impact the activity of only certain members of the enzyme family. In this paper, the catalytic mechanism of oxidation of DHO to OA in human DHOD was studied using a hybrid Quantum Mechanical/Molecular Mechanical (QM/MM) approach and Molecular Dynamics (MD) simulations. The free energy barriers calculated reveal that the mechanism in human DHOD occurs via a stepwise reaction pathway. In the first step, a proton is abstracted from the C5 of DHO to the deprotonated Ser215 side chain. Whereas, in the second step, the transfer of the hydride or hydride equivalent from the C6 of DHO to the N5 of FMN, where free energy barrier calculated by DFT/MM level is 10.84 kcal mol⁻¹. Finally, a residual decomposition analysis was carried out in order to elucidate the influence of the catalytic region residues during the DHO oxidation.

Introduction

Dihydroorotate dehydrogenases (DHODs)1 are flavin-containing enzymes that catalyze the fourth step in the de novo synthesis of pyrimidines. In this reaction, DHODs catalyze the conversion of dihydroorotate (DHO) to orotate (OA), the single redox step in pyrimidine synthesis. The diversity among DHODs of different organisms2 could enable the development of compounds that selectively inhibit de novo pyrimidine biosynthesis in some organisms while not affecting others. Therefore, DHOD enzymes represent an attractive and selective target for treating cancer, malaria, gastric ulcers, and rheumatoid arthritis.3–6 Many inhibitors have been developed to combat these diseases; some have been approved for use, as the case of the immunosuppressive drug leflunomide, which inhibits human DHOD and is used for the treatment of rheumatoid arthritis.5,7,8

DHODs can be classified into two classes based on sequence:1 class 1 and class 2. Class 1 are cytoplasmic enzymes and can be further subdivided into subclasses 1A and 1B.9,10 Class 1A enzymes (DHODs) form homodimers and appear to utilize Fumarate (FUM) as a physiological oxidant, in conjunction with oxidation of the reduced Flavin mononucleotide (FMN) cofactor during the second half-reaction.10 Family 1B forms heterotetramers and utilize NAD⁺ via a distinct protein subunit that contains a 2Fe–2S cluster and a Flavin Adenine dinucleotide (FAD) cofactor.9 Class 2 can be either homodimers or monomers and are membrane-bound enzymes that utilize respiratory quinone as a physiological oxidant during the second half-reaction.11

The mechanism of the dehydrogenation of dihydroorotic acid by DHOD has been the subject of extensive studies2,9,10,12–20. The oxidation reaction of DHO breaks two carbon-hydrogen bonds. An active site base (serine in class 2 enzymes or cysteine in class 1 enzymes) deprotonates C3 of DHO, and a hydride (or hydride equivalent) is transferred from C6 of DHO to N5 of the isoalloxazine ring of the flavin. The class 1 DHODs was demonstrated to follow a concerted mechanism.13–17 On the other hand, the studies by Fagan and coworkers, using DHO deuterated at the 5-, 6-, and both positions in anaerobic stopped-flow experiments, indicated that the class 2 enzyme from E. coli and H. sapiens use a stepwise mechanism.16,18 They suggested that when a stepwise mechanism occurs, there are two possible intermediates that could form: (i) if deprotonation occurs first, then an enolate intermediate will form, (ii) if hydride transfer occurs first, then an iminium intermediate will form (Scheme 1). It was not clear, in that study, which intermediate was forming in the stepwise oxidation of DHO by the class 2 DHOD enzymes.
Scheme 1 The proposed mechanisms for the DHO oxidation by human DHOD enzyme.

We present a computational free energy study aimed to elucidate which intermediate is formed in human DHOD a class 2 of DHOD enzymes. Here, a hybrid Quantum Mechanic/Molecular Mechanic (QM/MM) approach and Molecular Dynamic (MD) simulations is applied to study the details of the catalytic mechanism of DHO oxidation in the class 2. Hybrid QM/MM approaches combined with MD simulations, have been employed for many studies involving protein–inhibitors interaction and catalytic mechanism enzymatic. In the QM/MM methodology, one part of the system is described by quantum mechanics and the rest by molecular mechanics using a classical force field. In this study, the density functional tight binding (DFTB) semiempirical method to describe the QM region of the system. The DFTB method not only has been shown to be an accurate level of theory for describing the energetic of chemical reactions, but it was also demonstrated in biological systems that the minimum energy path results are in good agreement with higher levels of theory like MP2. A previous theoretical study also shown that DFTB method provides the best semiempirical description of six-membered carbohydrate ring deformation. Besides, recently, this semiempirical method was successfully applied to perform the catalytic mechanism for the DHOD from L. lactis, a class 2 DHOD enzyme. An umbrella sampling method using an adequate reaction coordinate was employed in order to obtain the free energy profile associated with the catalytic mechanism of DHO oxidation by human DHOD. In addition, a detailed analysis of the stabilization pattern of the active site residues on the intermediate (IT) and transition state (TS), in relation to the Michaelis Complex (MC), is examined through of energy decomposition methods.

Materials and methods

Initial system aspects and QM/MM simulations

The initial coordinates of human DHOD were obtained from the PDB database (access code: 1D3G, resolved at 1.60 Å). The human DHOD complex consist of 313 amino acid residues, 248 crystalline water molecules and the OA product and FMN cofactor bound in active site. In our simulations, the OA was “mutated”, in silico, to the DHO substrate. The missing hydrogen atoms were added to the system using the Leap module of Amber12 package taking into account a previous assignment of the protonation states of all these residues at pH 7 performed by recalculating the standard pKa values of the titratable amino acids using the empirical propKa program of Jensen et al. Particularly, the catalytic residue Ser215 was already considered in its active state (deprotonated), in accordance with experimental kinetics data. The system was immersed in a truncated octahedral cell of TIP3P water molecules, extending 10 Å outside the protein on each side. The force field parameters for the protein were assigned from the ff99SB, while the DHO substrate were obtained from gaff. The RESP approach was applied to obtain the substrate charges using HF/6-31G* level of theory calculated in Gaussian09 program. The FMN cofactor was described using parameter sets available in Bryce Group website.

The initial complex prepared as described on the previous paragraph were first submitted to 10000 steps of conjugate gradient energy minimization using the sander module of Amber12 package. Then, the complex was energy−minimized and then gradually heated to 300 K over 2 ns with a harmonic constraint value of 10 kcal·mol⁻¹·Å⁻² applied to the solute. Finally, the heated complex was equilibrated for 200 ps before the production stage. The minimizations, heating, equilibration and production stages employed a non-bonded cut-off of 8 Å. The particle mesh Ewald (PME) approach was applied to
calculate the long-range Coulomb Forces. The time step was set to 2 fs and all bonds involving hydrogen atoms were constrained using the SHAKE algorithm as during the MD simulation.

For the hybrid QM/MM simulations with DHOD-DHO complex, the atoms of the DHO, FMN and side chain of the deprotonated Ser215 amino acid residue were selected for QM region, which contains 49 atoms. The self-consistent charge density functional tight binding (sc-DFTB) semiempirical method, as implemented in Amber12, was selected to describe the QM region, while MM region of the complex (protein and water molecules) was described using the ff99SB and TIP3P parameter sets, respectively, as described early. The link atom method was used to complete the valence of the QM fragments into the QM-MM boundary in all part of QM system.

In order to analyze how the active site residues stabilize/desstabilize the INT and TS states in the catalyzed reaction of DHO by human DHOD with respect to MC state, we performed an energy decomposition analysis. This type of analysis has been extensively applied to enzymatic systems as the method has been discussed in detail elsewhere, here, just the main equations are presented.

\[ \Delta E_i = \Delta E_{i,\text{INT/TS}} - \Delta E_{i,\text{MC}} \] (2)

In our analysis, the QM subsystem was composed of the substrate, cofactor and Ser215. To obtain the average values for the interaction energy and stabilization effects of each residue in a particular state, 400 snapshots from QM/MM MD simulations were considered taken from umbrella sampling calculation with reaction coordinates corresponding to the MC, INT and TS. As no MD simulation is run when the i-th residue is mutated to Gly, the \( \Delta E_i \) does not consider the dynamic effects arising from differences in the structure/dynamics of the enzyme upon replacement. The \( \langle \Delta AE_i \rangle \) values provide hints about detrimental/beneficial effects of the i-th residue for the mechanism step under observation. Finally, these values cannot be considered to quantitatively estimate changes in the catalytic constant produced by mutation of the i-th residue by Gly for such changes depend on variations in the activation free energy, \( \Delta G^\ddagger \).

### Results and discussion

In the crystal structures of class 2 DHODs, residue Ser215 has been identified as the catalytic base, with one water molecule forming a bridge between Ser215 and Thr218 residues. In this enzyme, Ser215 is not exposed to solvent but a hydrogen bond network connects the active site base to bulk solvent. Small and coworkers using MD found the same hydrogen bond network which was observed experimentally. In that study, a proton relay mechanism had been hypothesized to facilitate the proton-transfer reaction from substrate to Ser215. In this sense, the proton would be passed from Cγ of DHO to the active site Ser215 to the water molecule in the tunnel, to a conserved Thr residue, and finally to bulk solvent, i.e., this network shuttles protons from the active site to bulk solvent and, therefore, could have a role in allowing the serine to act as a base. Fagan and co-workers through experimental evidences suggest a deprotonated state for the catalytic Ser residue. Besides, a recent computational study for the DHOD from *L. lactis* started the catalytic base (Cys residue) for the mechanism already in its deprotonated state. Moreover, DHOD enzymes from Class 1A have their active sites much more...
exposed to solvent than Class 2. Then, in these enzymes (Class 2), just an active serine alone is not enough; in this way, a fully proton relay network should be ordered for the most effective DHO deprotonation\textsuperscript{38}. Therefore, in our study the catalytic Ser215 residue was already considered in its deprotonated form given previous experimental and theoretical data\textsuperscript{11, 16, 18, 38, 59}. 

Umbrella Sampling and 2D-FES

In order to investigate the molecular mechanism of the oxidation of DHO to OA in human DHOD, we have employed umbrella sampling using DFTB/MM level. Figure 1 depicts the 2D-FES obtained through the RC1 \textit{versus} RC2 described in the previous section. The changes in the distances involved in the reaction coordinate definition, as well as averaged geometrical parameters for MC, INT, TS and P structures are presented in Table 1. As shown in Figure 1 and Table 1, the mechanism in human DHOD occurs via a stepwise reaction pathway. Importantly, this result supports the stepwise mechanism found experimentally for class 2 enzyme from \textit{E. coli} and \textit{H. sapiens}\textsuperscript{16, 18}. The first step of the molecular mechanism is the proton is abstracted from C\textsubscript{5} of DHO to deprotonated Ser215 side chain, and, the enolate intermediate is generated. In the second step the hydride or hydride equivalent is abstracted from C\textsubscript{6} of DHO to N\textsubscript{5} of FMN, completing the oxidation of the DHO into OA.

![Fig. 1 2D plot of FES for the DHO oxidation in human DHOD obtained by umbrella sampling simulations. The energy values are reported in kcal·mol\(^{-1}\)](image)

Table 1 Average distances collected from QM/MM MD simulations of reaction chemical species. The standard deviation values are present in parenthesis

<table>
<thead>
<tr>
<th></th>
<th>DFTB/MM</th>
<th>MC</th>
<th>INT</th>
<th>TS</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>d(C\textsubscript{5}-H\textsubscript{5})</td>
<td>1.25 (0.03)</td>
<td>2.30 (0.04)</td>
<td>2.30 (0.04)</td>
<td>2.29 (0.04)</td>
<td></td>
</tr>
<tr>
<td>d(H\textsubscript{5}-O\textsubscript{δ})</td>
<td>1.55 (0.04)</td>
<td>0.99 (0.02)</td>
<td>0.99 (0.03)</td>
<td>0.99 (0.03)</td>
<td></td>
</tr>
<tr>
<td>d(C\textsubscript{6}-H\textsubscript{6})</td>
<td>1.16 (0.04)</td>
<td>1.16 (0.03)</td>
<td>1.28 (0.04)</td>
<td>2.52 (0.04)</td>
<td></td>
</tr>
<tr>
<td>d(H\textsubscript{6}-N\textsubscript{5})</td>
<td>2.06 (0.04)</td>
<td>1.95 (0.04)</td>
<td>1.58 (0.04)</td>
<td>1.02 (0.03)</td>
<td></td>
</tr>
</tbody>
</table>

In the first step (proton abstraction step), the \(d_1\) bond distance changes from 1.25 Å in MC to 2.30 Å in INT, while the \(d_2\) distance changes from 1.55 Å in MC to 0.99 Å in INT, showing that proton transfer takes place before the hydride transference. Interestingly, during the proton abstraction step the transition state was not identified; an experimental kinetic isotope effects study for the class 2 enzyme from \textit{E. coli} show the deprotonation of C5 atom of DHO is fast enough that it does not contribute in the kinetic of the reaction\textsuperscript{16}. The computed \(\Delta G^\ddagger\) for this step is \(-6.54\) kcal·mol\(^{-1}\) making the INT more stable than MC, which agrees with previous experimental analyzes\textsuperscript{16}. In the second transfer of the hydride or hydride equivalent step, the \(d_3\) bond distance increases from 1.16 Å in INT to 1.28 Å in transition state (TS1) and then to 2.52 Å in P, while the \(d_4\) distance changes from
1.95 Å in INT to 1.58 Å in TS1 and then to 1.02 Å in P, indicating the transference of the hydride from C6 of DHO to N5 of FMN takes place after the proton abstraction step. On this step, the computed ∆G‡ is -13.50 kcal·mol⁻¹ making the transfer of the hydride thermodynamically favourable.

The free energy barrier (∆G‡) obtained by DFTB/MM level, between INT and P is 2.04 kcal·mol⁻¹ (10.84 and 7.55 kcal·mol⁻¹ using B3LYP-D3/MM and MP2/MM corrections, respectively) for the hydride equivalent step. This result rationalize the experimental evidence that identifies DHOD as a very fast enzyme (fast enough that the experimental procedures must be carried out at low temperatures to be measurable) and suggests that DHO oxidation occurs via a stepwise mechanism for DHOD class 2 enzymes, which is in agreement with mechanisms proposed by Fagan and coworkers.[16]

Finally, in order to evaluate the mechanism found at DFTB/MM level, we also performed B3LYP-D3/MM and MP2/MM calculations. In this calculation, we carried out single-point calculations using representative snapshots from MC, INT, TS and P states. It confirms that the proton abstraction step is kinetic and thermodynamically followed by the hydride equivalent step (see supplementary data).

**Interaction Energy Decomposition**

The origin of the substrate-enzyme interaction can be analyzed by decomposing the total interaction energy and taking into account the difference of energies when a particular residue is present, as described in the method section. In order to get a deep insight into protein-substrate interactions, 400 snapshots from QM/MM MD simulations were considered taken from umbrella sampling calculation, for substrate-protein complexes appearing along the 2D-FES (MC, INT and TS). Figure 2a shows how the main amino acid residues interact in the active site. Averaged protein-substrate interaction energies by residue in MC and relative stabilization pattern on the INT and TS with respect to the MC are displayed in Figure 2b. The addition over all the ∆AE values for each residue represents an indication of the degree of the stabilization of the INT and TS from the enzyme environment on the QM subsystem. The resultant values were -19.03 and -7.12 kcal·mol⁻¹, indicating a strong stabilizing effect on the INT and TS, respectively. From Figure 2a we can observe how the MC interacts through hydrogen bonds, with a pocket created by Lys100, Asn145, Asn212, Asn217, Asn284 and Thr285 residues. The Asn145, Asn212, Asn217, Asn284 and Thr285 residues are conserved in all DHODs sequences[25, 60-65]. Lys100, which is conserved in family 2 members, also form also hydrogen bond with FMN. The specific function of each residue during the DHO oxidation mechanism is still unknown[15]. They would play a role in the conversion of the DHO into OA by stabilizing the chemical species found during the catalytic mechanism, i.e., the transition state and/or the intermediate, as well as the binding of DHO for the reaction. Then, in order to elucidate and to try better understanding their roles, an interaction energy decomposition analysis was computed. In Figure 2b, positive values correspond to unfavorable interactions, whereas negative values mean that the interaction is stabilizing the substrate in that particular state.

The Figure 2b shows the relative stabilization pattern on the INT and TS with respect to the MC. The INT enzyme complex presents a stabilization pattern stronger to the one observed for TS enzyme. Thus, our analysis will be focused on the INT and TS. A detailed analysis of the stabilization pattern residues to the total INT complex and TS complex stabilization energy shows that while the strongest interaction of INT is established with the Lys100, Asn145, Asn212, Asn217, Asn284 and Thr285 residues, while in the case of TS2, it is with just Lys100, Asn217 and Asn284 residues. The Asn212 and Asn284 are well positioned to stabilize any partial negative charge that developed at the DHO in INT and, therefore, it stabilizes the formation of enolate intermediate, while Asn154 does not show an important role during stabilizing any chemical species, once that iminium intermediate is not created during the DHO oxidation. Therefore, these results evidence the formation of enolate as the reaction intermediate. In fact, Fagan and co-workers demonstrated that the mutation of Asn212 and Asn284 results in a large decrease in the reduction rate constant (10500-fold and 2000-fold, respectively), i.e., the Asn212 and Asn284 residues are important for catalytic

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**Fig. 2 (A)** The representative MC frame on 2D-FES considered on the residual decomposition analysis. The QM subsystem is presented on green carbon atoms. **(B)** Plot of stabilization of the TS (red) and INT (blue) states with respect to the MC by active region residues.
Conclusions

We have investigated the mechanism of the catalytic mechanism of oxidation of DHO to OA in human DHOD through DFTB/MM MD calculations. The results obtained show that the mechanism in human DHOD occurs via a stepwise reaction pathway. In the first step of the molecular mechanism the proton is abstracted from C$_5$ of DHO to deprotonated Ser215 side chain, and, in the second step the transfer of the hydride or hydride equivalent from C$_6$ of DHO to N$_8$ of FMN occurs. The $\Delta G^i$ obtained for the reaction by DFTB/MM level is 2.04 kcal·mol$^{-1}$ for second step; however, by means B3LYP-D3/MM and MP2/MM single point calculations, the $\Delta G^i$ value increase to 10.84 and 7.55 kcal·mol$^{-1}$, respectively, which is suitable to experimental evidences. As observed in the analysis of individual interactions, the influence of Lys100, Asn212 and Asn284 seems to be crucial for FMN reduction. Thus, the present theoretical study is not only important to get an insight into the reaction, but the knowledge of the molecular mechanism can be used to get structural and electronic information of the species appearing along an enzymatic reaction path, which is of great value for a systematic protocol for synthesis of inhibitors. We hope it can contribute to future studies of this important system and could eventually facilitate development of the inhibitors for treating cancer, malaria, gastric ulcers, and rheumatoid arthritis.

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Notes

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$^c$ Electronic Supplementary Information (ESI) available: [details of the free energy barrier correction to the SCC-DFTB/MM by B3LYP-D3/MM and MP2/MM]. See DOI: 10.1039/b000000x/

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