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Binding mechanism of uranyl to transferrin implicated by density functional theory study†

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Transferrins have been proposed to be responsible for the *in vivo* transportation of uranyl. In this work, the binding mechanism of uranyl to transferrin has been studied using density functional theory method. Three possible stepwise pathways have been investigated and compared, differing in the sequence of the three residues to bind with uranyl, *i.e.* $Tyr^* \rightarrow Tyr^* \rightarrow Asp^*$ (YYD) and $Tyr^* \rightarrow Asp^* \rightarrow Tyr^*$ (YDY) and $Asp^* \rightarrow Tyr^*$ (DYY). Compared with the activation energies and the reaction heat of these three possible mechanisms, it is concluded that the YYD pathway is a more plausible description for the binding of uranyl. According to the calculations, the binding process is described as a ligand exchange process assisted by the hydrolysis of uranyl tricarbonate complex, and the role of carbonate ligand which determines the optimal pathway is identified. The QTAIM analysis was used to compare the bond nature of uranyl complexes in its free form and its complex with the amino acid residues. The results are expected to benefit our understanding of the uptake of uranyl by serum transferrins, and have implications on protein engineering and the development of decorporation agents towards improved binding kinetics and thermodynamics of uranyl in a specific pH range.

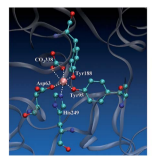
Introduction

Uranium is a toxic and radioactive heavy metal. It is naturally available in the earth's crust, and can be found in, *e.g.* food, groundwater, and mineral resources. Uranium distributed in the environment may enter the human body *via* various ways. *In vivo* uranium is toxic to bones, kidneys, lungs, and the brain, ²⁻⁶ and can even cause cancer in the human body. In serum, uranium predominantly exists in the uranyl $(UO_2^{\ 2^+})$ form, ⁷⁻¹⁰ and displays an affinity constant in the range of 10^7 to 10^{13} to combine with serum transferrin (sTf), ^{9,11,12} and the uranyl–sTf complex can form a weak interaction with the sTf receptor to

constitute a potential channel for the transportation of uranyl from serum to cell.¹³

Serum transferrin is a glycoprotein of about 80 kDa, and its major function is to transport Fe³⁺ in human cells.¹⁴ It exists in the form of a single chain with two lobes, C and N lobes, consisting of 700 amino acid residues. Each lobe has an ironbinding site constituted by four residues, including two tyrosines (Tyr), one histidine (His), and one aspartate (Asp).¹⁵ The N-lobe of transferrin in its holo state has been crystallized¹⁶ and its binding site with ferric cation bound is shown in Fig. 1 to highlight how the key residues assembly to accommodate the ferric cation. When the transferrin is in its apo state, the four ligands open up in the aqueous medium.¹⁷⁻¹⁹ The apo state can

 $[\]dagger$ Electronic supplementary information (ESI) available: The full citation of Gaussian 09; discussion on channel III of the first HTyr* binding to UO2(CO3)3 $^{4-}$ along pathway YYD; the energy, entropy, and free energy of each stationary point involved in eqn (1)–(7) and the relative energy, entropy, and free energy of eqn (1)–(7); the schematic structures of stationary points involved in the present work while not included in the main text along with key geometrical parameters; the natural charges of key atoms and the Mayer atomic bond order of key bond along the reaction coordination of Tyr* binding toward uranyl; the coordinates and energies of stationary points. See DOI: 10.1039/c6ra26109d



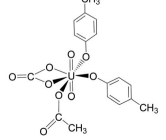


Fig. 1 The binding site of the N-lobe of the human serum transferrin with $\mathrm{Fe^{3+}}$ bound (PDB ID $\mathrm{1a8e^{16}}$) (left) and the schematic elucidation of its possible binding conformation with uranyl considered in the present work (right), where the Tyr and Asp residues are simplified as p-cresol and acetate, respectively.

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also form complex with other metals besides iron, such as aluminum, titanium, bismuth, gallium, cobalt, uranium, and so on. $^{20-22}$

Uranyl differs from Fe3+ in its linear geometry with two Oyl atoms at the axial coordination sites. When coordinating with sTf, since the axial positions of uranyl are occupied by O^{yl} groups only three ligands may bind with uranyl in the equatorial plane in addition to a synergistic bidentate ligand, and the binding region of the complex is semi-closed.10 In a previous study, Vidaud et al. obtained the FTIR spectra of the uranyltransferrin complex, which showed the His249 does not connect to the uranyl ion.23 This was later supported by a B3LYP study of the vibrational frequencies of potential QM models.24 The stronger interaction between O and U than that between N and U was also observed in a survey of crystal structures of proteins with uranyl bound deposited in the databases.25 In the work of Vidaud et al.,23 the binding of uranyl has been compared to that of Fe³⁺, and a structural model was proposed to illustrate the semi-open conformation of uranyl-sTf complex which may be responsible to its weaker interaction with the sTf receptor than that of Fe³⁺-sTf. In that work, the Asp63 and the synergistic anion CO₃²⁻ were identified to interact with uranyl differently compared to Fe³⁺, while lack of details at the molecular level in their interaction modes with uranyl.

The burgeoning application of fission energy in civilian use raises serious concerns on the environmental and health consequences. The insufficient knowledge on the in vivo migration mechanism of uranyl leaves the public in fear on the operation of nuclear plant, but also makes it difficult to implement a rescue strategy to cope with unexpected situations. This motivated the current work, in which we employed density functional theory method to investigate the binding process of uranyl to sTf in serum. Based on previous work from both experimental15 and computational24,26,27 sides, we assumed a stepwise pathway started from uranyl tricarbonate, during which the U atom remains in hexa- or penta-coordinated. This work is expected to shed light on our understanding of the in vivo behavior of uranyl and has implications on protein engineering targeting at improved binding kinetics and thermodynamics of uranyl and biomolecules or at the development of decorporation agents.

Computational details

Density functional theory (DFT) methods have been used²⁸⁻³¹ in the studies of actinide chemistry. In this work, B3LYP functional,³²⁻³⁵ as implemented in Gaussian 09 package,³⁶ was employed to conduct the calculations. Uranium was described by a quasi-relativistic pseudopotential according to Dolg, et al.,³⁷⁻³⁹ denoted as ECP60MWB, which treats inner 60 electrons in the core shells, together with the optimized basis set for the valence shells. For the other atoms, a triple- ζ -quality basis set with diffuse and polarization functions, denoted as 6-311++G(d,p), was used. All stationary points were fully optimized, followed by vibrational frequency calculations to identify the nature of each structure to be a minimum or a transition state. Intrinsic reaction coordinate (IRC)⁴⁰ calculations were

carried out to guarantee that the transition states are in the right reaction paths. Correction to the Gibbs free energy of solvation was done by a THERMO program according to Fang.⁴¹ All discussions about the thermodynamics of the reaction pathways are based on the corrected Gibbs free energies.

Simplified models cultivated from the binding site of sTf were used to study the binding mechanism of uranyl. Only the functional groups of the residues that are responsible for the binding of uranyl to sTf were considered, and the solvent environment was handled by using the polarizable continuum model (PCM)^{42,43} with a dielectric permittivity of 78.3553 (water). In the models, the Tyr was mimicked by *p*-cresol (HTyr*) and Asp by acetate (HAsp*). The protonation states of the residues were determined according to their pK_a values, which are 9.84 \pm 0.11 for Tyr and 3.67 \pm 0.04 for Asp, respectively,⁴⁴ thus at pH = 7, the Tyr exists in its protonated state while Asp in its deprotonated state.

Results and discussion

Previous studies^{45,46} show that the *in vivo* uranyl may exist in the free form, or binding with target proteins, *e.g.* transferrins, or deposited in bones. The general form of free uranyl in serum appears as uranyl tricarbonate complex $(UO_2(CO_3)_3^{4-})$, which was used as the starting material in the present study. In the holo form of transferrins, the uranyl interacts with the protein *via* the coordination with three amino acid residues, *i.e.* two tyrosine and one aspartic acid. We consider the binding process of one lobe of transferrins. Assuming the binding of uranyl to the transferrin proceeds in a stepwise manner, three pathways were investigated, differing in the sequence of the residues to coordinate to uranyl: $Tyr^* \rightarrow Tyr^* \rightarrow Asp^*$ (denoted as YYD), $Tyr^* \rightarrow Asp^* \rightarrow Tyr^*$ (YDY) or $Asp^* \rightarrow Tyr^* \rightarrow Tyr^*$ (DYY). In the following, these three pathways will be analyzed and compared.

I. Pathway YYD: $Tyr^* \rightarrow Tyr^* \rightarrow Asp^*$

Along this pathway, the whole reaction starts two sequential ligand exchange steps of carbonate by HTyr*. The released excess coordination site of uranyl is then occupied by an Asp* residue nearby. Initially, the exchange of carbonate by the first Tyr* may happen in three channels, *i.e.* either by a direct ligand exchange between the two groups (channel I, eqn (1)), or assisted by a hydronium ion in aqueous phase (channel II, eqn (2) and (3)), or a direct ligand exchange between hydrolyzed uranyl tricarbonate species and Tyr* (channel III, eqn (4) and (5)). These three channels are elucidated as below:

Channel I:

$$UO_2(CO_3)_3^{4-}$$
 (1a) + HTyr* \rightarrow $UO_2(CO_3)_2(Tyr^*)^{3-}$ (1b) + HCO₃⁻ (1)

Channel II:

$$UO_2(CO_3)_3^{4-}$$
 (1a) + $H_3O^+ \rightarrow UO_2(CO_3)_2(H_2O)^{2-}$ (U- H_2O) + HCO_3^- (2)

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$$\begin{array}{c} UO_{2}(CO_{3})_{2}(H_{2}O)^{2-}\left(U\text{-}H_{2}O\right)+HTyr^{*}\rightarrow \\ UO_{2}(CO_{3})_{2}(Tyr^{*})^{3-}\left(1b\right)+H_{3}O^{+} \end{array} \tag{3}$$

Channel III:

$$UO_2(CO_3)_3^{4-}$$
 (1a) + H₂O \rightarrow $UO_2(CO_3)_2(OH)^{3-}$ (U-OH) + HCO₃⁻ (4)

$$UO_2(CO_3)_2(OH)^{3-} (U-OH) + HTyr^* \rightarrow UO_2(CO_3)_2(Tyr^*)^{3-} (1b) + H_2O$$
 (5)

According to the calculations, the direct ligand exchange reaction (channel I) is an exothermic process with a free energy change of -8.0 kcal mol⁻¹ (Table S1†). The channel II mimics an acidic environment and the hydration of uranyl tricarbonate complex 1a is strongly exothermic by 53.2 kcal mol^{-1} . However, the highly stable hydrated intermediate UO₂(CO₃)₂(H₂O)²⁻ (U-H₂O) is inert to the ligand exchange of H₂O by HTyr* thermodynamically and the reaction is endothermic by $45.2 \text{ kcal mol}^{-1}$. This suggests that in acidic environment, the uranyl species may be trapped in its UO₂(CO₃)₂(H₂O)²⁻ state, which hammers its binding with the HTyr* ligand. The channel III is a ligand exchange reaction assisted by the participation of water. The reaction of eqn (4) is moderately endothermic by $2.2 \text{ kcal mol}^{-1}$. The HTyr* then approaches the uranyl to replace the hydroxide anion (eqn (5)). This reaction is thermodynamically favorable in view of its exothermicity of -10.2 kcal mol⁻¹. Based on the above data, below we consider the reaction initialized by the channel I. The channel III was also investigated and the data are provided in ESI.†

Along the channel I, starting from 1a and HTyr*, the first ligand exchange reaction is described as a four-step process, including the breakage of U-O4 bond (TS1a), the proton transfer from HTyr* to O5 (TS2a), the ligand exchange of HCO₃ by $Tyr^*(TS3a)$, and the HCO_3^- release (TS4a), as shown in Fig. 2. The free energy profile is plotted in Fig. 3. When approaching 1a, HTyr* may first form a hydrogen bond (H-bond) with one carbonate group (COM1a), which is thermodynamically favorable by 7.4 kcal mol⁻¹. In COM1a, the HTyr* interacts with the carbonate group via an H-bond between H6 (HTyr*) and O5 (carbonate). We have also checked the possibility for an H-bond between H6 and O2/O4, which is given in Fig. S4,† and found that the complex is less stable than COM1a by about 1.6 kcal mol^{-1} . This is because O5 (-0.84e) is more negatively charged than O2/O4 (-0.74e/-0.74e), leading to stronger interaction between H6 and O5 than that between H6 and O2/O4. Below we consider the COM1a as a key species on the pathway of the ligand exchange reaction.

The H-bond interaction between HTyr* and CO₃²⁻ may perturb the dative bond between the carbonate group and uranyl, and induce a transition of the coordination mode of CO_3^{2-} from κ^2 in COM1a to η^1 in INT1a via a transition state TS1a by overcoming a free energy barrier of 4.9 kcal mol⁻¹. The highly nucleophilic CO₃²⁻ group may abstract the proton H6 from the Tyr* group, as indicated by the variation of the Mayer atomic bond orders (MBO) of H6-O5, which is to be formed, and H6-O7, which is to be broken, from 0.123 and 0.568 in

INT1a to 0.378 and 0.235 in TS2a to 0.515 and 0.177 in INT2a, respectively (Fig. 4b).

The protonated carbonate group is easily replaced by the Tyr* anion via TS3a. In TS3a, the imaginary vibration frequency corresponds to the asymmetric stretching mode of the U-O7 (Tyr*) and U-O2 (carbonate), suggesting the coordination of Tyr* and the dissociation of bicarbonate happened simultaneously. The free energy barrier to this process is 13.1 kcal mol⁻¹ and this step is endothermic by 9.4 kcal mol⁻¹. The detached HCO₃⁻ group then leaves, and gives the penta-coordinated UO₂(CO₃)₂ $(Tyr^*)^{3-}$ complex (INT4a). In INT4a, the Tyr* is coordinated to U in a η¹-manner, with a U-O7 bond length of 2.306 Å. Note that, the coordination of the second deprotonated phenolate ligand (INT2b \rightarrow TS3b \rightarrow INT3b) is exothermic, which is in contrast to that of the first one (INT2a \rightarrow TS3a \rightarrow INT3a). This is because that INT2b is less negatively charged than INT2a. In the meantime, uranyl has coordination site ready to receive the deprotonated phenolate ligand. This leads to a thermodynamically more favorable formation of INT3b than INT3a.

In Fig. 4, the fluctuations of the natural charges of U and the coordinating O atoms of the ligands and the MBO of key bonds along the channel I are monitored, which are consistent with the structural evolution along the reaction. Considering the proton transfer step as an example, it can be seen that accompanying the proton transfer from HTyr* (O7) to one carbonate ligand (O5), there is marginal change in the atomic charges residing on these two atoms and the O4 atom. For the other atoms, the atomic charges remain roughly constant during the reaction. This suggests that the complex may adapt to the new chemical environment during the ligand exchange and reach its equilibrium state quickly. The monitoring of the MBO clearly shows the cleavage of U1-O2, U1-O4, and O7-H6 bonds, whose MBO values evolve from 0.4-0.6 to around 0.0, and the formation of O5-H6 and U1-O7 bonds.

The newly formed INT4a may experience a second ligand exchange reaction to have one of the rest two carbonate groups replaced by another HTyr* ligand. A process similar to the first ligand exchange reaction was identified, as shown in Fig. 2.

Similar to the binding of the first Tyr*, along the binding of the second Tyr* ligand, there is also no significant charge transfer (within 0.28e) during the reaction, as seen in Fig. 4c, and the evolution of the Mayer atomic bond order of key bonds (Fig. 4d) clearly shows that the proton transfer from O14 (HTyr*) to O12 (CO_3^{2-}) happens prior to the U-O14 bond formation and the U-O11 bond cleavage.

Upon the dissociation of O14···H13-O12 hydrogen bond in INT4b, a water molecule close to uranyl may attach to U to keep it penta-coordinated in its equatorial plane. This gives the intermediate $UO_2(CO_3)(H_2O)(Tyr^*)_2^{2-}$ (1c, Fig. 2). It may interact to the neighboring Asp carboxylate group, which is modelled by Asp* ligand here. The reaction is a thermoneutral process with the free energy of Prod being 1.1 kcal mol⁻¹ lower than 1c. The free energy profile of this whole process of YYD is plotted in Fig. 3.

In summary, following the pathway YYD starting with channel I, the rate-determining step is the coordination of the first Tyr* ligand (TS3a) with a relative free energy of 8.2 kcal RSC Advances Paper

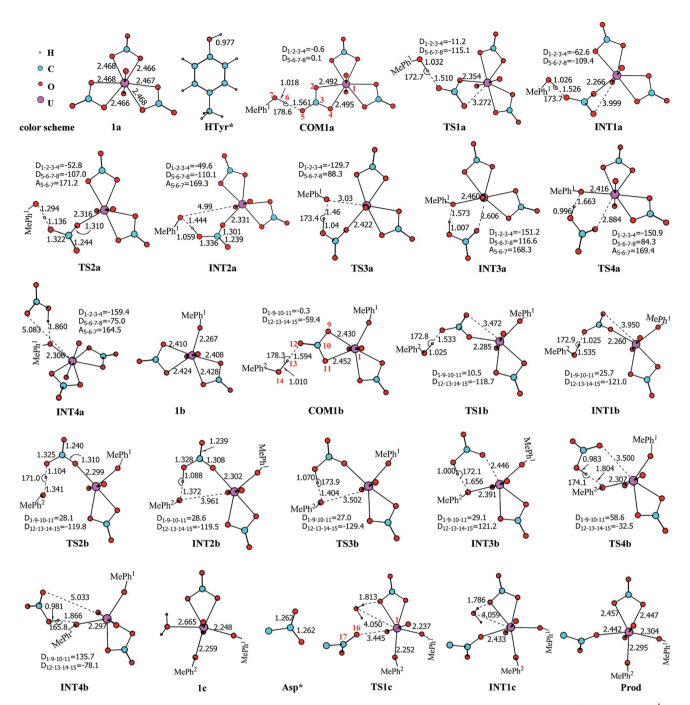


Fig. 2 The schematic structures of stationary points following the YYD pathway along with key geometrical parameters (bond lengths in Å, and bond angles and dihedral angles in degree). The numbering of key atoms is given in COM1a, COM1b, and TS1c, and the aromatic C atom of p-cresol that is bonded to its hydroxyl group is numbered as 8 in the first HTyr* ligand or 15 in the second one. The aliphatic hydrogen atoms of Asp* are omitted for simplicity. The solid lines represent the normal covalent or coordination bonds, and the dash lines represent the partially cleaved/formed bonds or weak interactions.

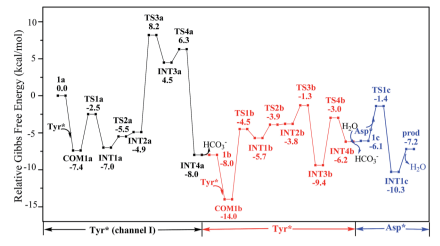
mol⁻¹, and the entire reaction is exothermic by 7.2 kcal mol⁻¹. This suggests that the whole reaction can happen easily under the physiological condition.

II. Pathway YDY: $Tyr^* \rightarrow Asp^* \rightarrow Tyr^*$

The pathway YDY differs from the YYD in the second ligand exchange step, where in the former case a negatively charged

Asp* ligand, instead of Tyr* in the latter case, reacts with $UO_2(CO_3)_2(Tyr^*)^{3-}$ (**1b**). Here starting from $UO_2(CO_3)_2(Tyr^*)^{3-}$, we located the stationary points of the second and the third ligand exchange steps in the pathway YDY. The optimized structures are shown in Fig. 5 for the two sequential steps, and the free energy profile of process YDY is plotted in Fig. 6.

For the coordination of Asp*, as the carboxylate group is in its deprotonated state, we did not consider the release of one



Gibbs free energy profile of the YYD pathway.

carbonate group upon its coordination, but considered a direct coordination to U atom at the site between Tyr* and a carbonate group. This process faces a high free energy barrier of up to 20.1 kcal mol⁻¹ (TS1b') and is strongly endothermic by 17.7 kcal mol⁻¹ (INT1b'). In the transition state TS1b', the interaction between U and the original three ligands is weakened in view of the longer U-O distances, and the U-O16 (Asp*) bond which is going to form is calculated to have a distance of 2.92 Å. The imaginary frequency is -84.2 cm⁻¹, corresponding to the vibrational mode of U-O16 (Asp*) stretching. The binding of Asp* gives a hexa-coordinated complex INT1b'.

As in INT1b' the U atom is hexa-coordinated in its equatorial plane, the binding of the third ligand Tyr* requires the release of at least one coordination site. Concerning the binding mode of uranyl in sTf, here we consider the replacement of one of the two carbonate groups by a second Tyr* ligand. Similar to the coordination of the first Tyr* ligand shown in Fig. 2, the HTyr* ligand first builds an H-bond interaction with the leaving CO_3^{2-} group. The hydrogen bond between O12 (carbonate) and H13 (Tyr*) stabilizes COM1c' by 5.7 kcal mol⁻¹. The following cleavage of the U-O9 coordination bond, which releases one coordination site, is a facile process with a free energy barrier of 2.1 kcal mol^{-1} (TS1c') and an exothermicity of 7.2 kcal mol^{-1} (INT1c'). The η^1 -coordinated carbonate group is highly nucleophilic, and can easily abstract a proton (H13) from the HTyr* (TS2c'). The formation of a dative bond between U and O14 (Tyr*) gives the product via TS3c', at the cost of the break of U-O11 (carbonate) bond with a free energy barrier of 10.7 kcal mol⁻¹. The whole reaction of the pathway YDY is exothermic by $7.2 \text{ kcal mol}^{-1}$.

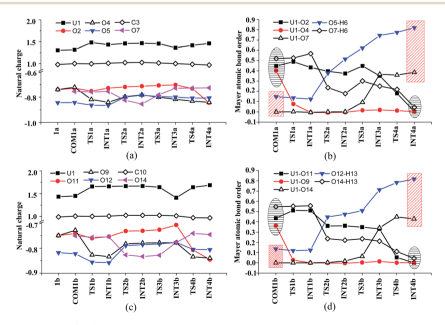


Fig. 4 The fluctuation of natural charge (in e) of key atoms and Mayer atomic bond order of key bonds during the ligand exchange of two CO₃²⁻ by the first HTyr* (a and b) and by the second HTyr* ligand (c and d).

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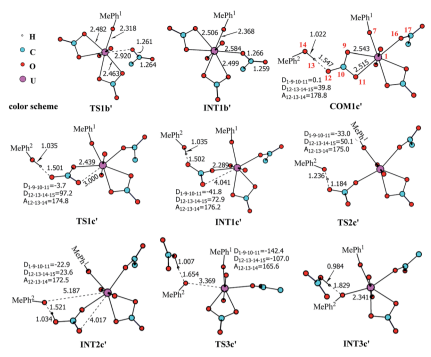


Fig. 5 The schematic structures of stationary points along the binding of Asp* to $UO_2(CO_3)_2(Tyr^*)^{3-}$ along with key geometrical parameters (bond lengths in Å, and bond angles and dihedral angles in degree). The numbering of selected atoms is shown in 1b and COM1c'. The aliphatic hydrogen atoms of Asp* are omitted for simplicity. The solid lines represent the normal covalent or coordination bonds, and the dash lines represent the partially cleaved/formed bonds or weak interactions.

III. Pathway DYY: $Asp^* \rightarrow Tyr^* \rightarrow Tyr^*$

In principle, there exists a third potential pathway for the binding of UO₂(CO₃)₃⁴⁻ to sTf in a sequential manner. Along this pathway, the uptake of uranyl by sTf starts from its coordination to a negatively charged Asp residue followed by the binding of two Tyr residues nearby. With the models employed here, this pathway is denoted as $Asp^* \rightarrow Tyr^* \rightarrow Tyr^*$ (DYY). We first considered the formation of the intermediate UO₂ $(CO_3)_3(Asp^*)^{5-}$ (U-3CO₃-Asp*, Fig. 7) via the reaction below:

$$UO_2(CO_3)_3^{4-}$$
 (1a) + Asp* \rightarrow $UO_2(CO_3)_3(Asp*)^{5-}$ (U-3CO₃-Asp*) (6)

Thermodynamically this reaction is unfavorable and the Gibbs free energy change was calculated to be 16.2 kcal mol⁻¹ (Table S1†). This means that an activation free energy more than 16.2 kcal mol⁻¹ is inherent to the binding of Asp* to the $UO_2(CO_3)_3^{4-}$ complex. It is conceivable that the negatively charged Asp* feels strong repulsive interaction when

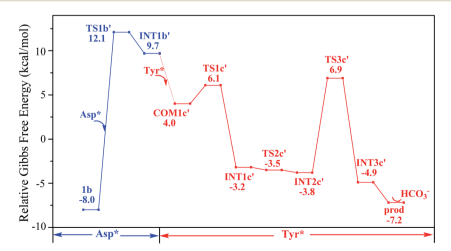


Fig. 6 Gibbs free energy profile of the sequential binding processes of Asp* (blue), and the second HTyr* (red) along the YDY pathway. The first HTyr* binding is the same as that in YYD pathway, which is not repeated here.

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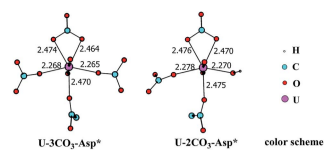


Fig. 7 The schematic structures of U-3CO₃-Asp* and U-2CO₃-Asp* with key geometrical parameters (bond lengths in Å).

approaching to the negatively charged $UO_2(CO_3)_3^{4-}$ complex which blocks this channel. This suggests that the direct exchange of CO_3^{2-} by Asp* is hard to happen.

There exists another possibility with the exchange mediated by a water molecule, which may thus alleviate the electrostatic repulsion. It involves two reactions below:

$$UO_2(CO_3)_3^{4-}$$
 (1a) + H₂O \rightarrow
 $UO_2(CO_3)_2(OH)^{3-}$ (U-OH) + HCO₃⁻ (4)

$$UO_2(CO_3)_2(OH)^{3-} (U-OH) + Asp^{*-} \rightarrow UO_2(CO_3)_2(OH)(Asp^*)^{4-} (U-2CO_3-Asp^*)$$
 (7)

This clears space in the surrounding of U atom to allow Asp* to approach uranyl. The ΔG of eqn (7) is calculated to be 11.3 kcal mol⁻¹. These data suggest that the DYY pathway cannot compete against the YYD pathway under physiological conditions.

IV. Discussion

We note that in an earlier B3LYP study,²⁴ an uranyl complex was constructed with the equatorial plane of U atom tetracoordinated, where the Asp and the carbonate groups coordinate with uranyl as monodentate ligands. In a well confined and limited space, such a conformation may represent a special case for uranyl to adapt to the chemical environment constructed by protein due to the constraints of protein backbone. In recent years, it is established knowledge that in aqueous phase, uranyl prefers to adopt a penta-coordinated conformation in its equatorial plane. In the case of sTf, the uptake of the uranyl triggers a domain rotation of the protein around an axis crossing the hinge site. This means that the binding site is not

uniquely defined prior to the binding of uranyl, and the flexible residues may adjust their orientations to reach an optimal binding state with uranyl. Thus, in this work, we have considered a hexa- or penta-coordinated conformation of the uranyl in its equatorial plane, where the synergistic anion binds to uranyl in a κ^2 manner. According to our calculations, the whole reaction is exothermic, suggesting such a treatment is reasonable in view of thermodynamics.

As mentioned above, the free energy barrier to the ratedetermining step of pathway YYD is 13.1 kcal mol⁻¹ which corresponds to the ligand exchange of the first Tyr* with HCO₃⁻. For the pathway YDY, the binding of the Asp* ligand costs more energy of 20.1 kcal mol⁻¹ (Asp*) and appears as the rate-determining step. These data show that when loading to sTf, the uranyl tricarbonate favors to coordinate to two Tyr residues sequentially first prior to its interaction with Asp, although Asp is ready to interact with uranyl while Tyr requires to be deprotonated before coordinating to uranyl. Note that, in serum, uranyl exists mainly in its tricarbonate complex form, and any ligand exchange process requires the release of one or two of its carbonate ligands. Keeping this in mind, the proton on the phenol group offers a "dragging force" to weaken the interaction between the coordinated carbonate ligand and uranyl and facilitate its leaving to trigger the ligand exchange reaction, in Chinese idiom, mislead a tiger (carbonate) to leave the mountains it based (uranyl) by a plot (proton) in order to conquer its territory (the coordination site). This benefits the Tyr residue to grab uranyl nearby. The deprotonated Asp residue does not have this "bait", and cannot compete against Tyr residue in their coordinating with uranyl which is under the protection of the negatively charged carbonate ligands. In other words, the approach of Asp to uranyl is posterior to the binding of two Tyr residues, and the YYD pathway is the most plausible mechanism for the ligand exchange of uranium tricarbonate. An overview of the YYD pathway is shown in Fig. 8.

According to our calculations, the whole process is exothermic, either along the YYD or the YDY pathway, by -7.2 kcal mol^{-1} . This suggests that the capture of uranyl by sTf is a thermodynamically favorable process. To understand the bonding nature of the starting material $\mathrm{UO_2(CO_3)_3}^{4-}$ and the product $\mathrm{UO_2(CO_3)(Tyr^*)_2(Asp^*)^{3-}}$, we analyzed the bonding properties of the coordination bonds using QTAIM topological analysis technique. Four descriptors are used to compare the coordination bonds in the two complexes, and these include the electronic density $\rho_{\rm b}$ and its Laplacian $\nabla^2 \rho_{\rm b}$, the energy density

Fig. 8 The most probable mechanism (YYD) for the binding of the uranyl to transferrin.

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Table 1 The $ρ_b$ (e bohr⁻³), $\nabla^2 ρ_b$ (e bohr⁻⁵), $H_b(r)$ (hartree bohr⁻³), and δ of the coordination bonds of UO₂(CO₃)₃⁴⁻ (1a) and UO₂(CO₃)(-Tyr*)₂(Asp*)³⁻ (prod) from QTAIM analysis

$UO_2(CO_3)_3^{4-}$					$UO_2(CO_3)(Tyr^*)_2(Asp^*)^{3-}$				
$Bond^a$	$ ho_{ m b}$	$\nabla^2 \rho_{\rm b}$	$H_{\rm b}(r)$	δ	$Bond^a$	$ ho_{ m b}$	$\nabla^2 \rho_{\rm b}$	$H_{\rm b}(r)$	δ
U-O ^{1a}	0.0592	0.1856	-0.0045	1.12	U-O ^{1a}	0.0623	0.1964	-0.0053	1.16
$U-O^{1b}$	0.0595	0.1863	-0.0046	1.12	$U-O^{1b}$	0.0609	0.1920	-0.0050	1.14
$U-O^{2a}$	0.0594	0.1863	-0.0046	1.12	$U-O^{Y}$	0.0763	0.3021	-0.0084	1.39
$U-O^{2b}$	0.0592	0.1858	-0.0045	1.12	$U-O^{Y'}$	0.0776	0.3070	-0.0090	1.42
$U-O^{3a}$	0.0593	0.1859	-0.0045	1.12	$U-O^D$	0.0557	0.2168	-0.0017	1.13
$U-O^{3b}$	0.0596	0.1867	-0.0046	1.12					

^a O^{ia} and O^{ib} (i = 1, 2, 3) represent the two coordinating O atoms of the *i*-th CO_3^{2-} ; O^Y , $O^{Y'}$ and O^D are the coordinating O atoms of the two Tyr* (Y, Y')and the Asp* (D) ligands, respectively.

 $H_{\rm b}(r)$ at the bond critical point, and the delocalization index between two atoms $\delta(A,B)$. These values are collected in Table 1.

As seen in Table 1, in $UO_2(CO_3)_3^{4-}$ (1a), the six U-O (carbonate) bonds have almost same nature with a $\rho_{\rm b}$ of 0.06 e bohr⁻³ and a $\nabla^2 \rho_b$ of 0.19 e bohr⁻⁵, suggesting a depleted feature for the interaction between the two atoms in each pair. Upon the substitution of two CO₃²⁻ ligands by two Tyr* and one Asp* ligands to form UO₂(CO₃)(Tyr*)₂(Asp*)³⁻ (prod), as the coordination number decreases from 6 to 5, the values of $\delta(A,B)$ for all of the five dative bonds increase marginally compared to those in the reactant with the largest values for U-O (Tyr*) bonds which are 1.39 and 1.42, respectively. We note that the U- η^{1} -O (Asp*) has a smaller ρ_{b} while a larger $\nabla^{2}\rho_{b}$ than the U- κ^{2} -O (carbonate), suggesting different bond ellipticity between them. The larger ρ_b of U–O (Tyr*) than the other coordination bonds indicates its more covalent feature due to its stronger nucleophilicity.

Conclusion

The binding process of uranium to transferrin has been investigated by density functional theory method, and three stepwise pathways have been compared, differing in the sequence of the three amino acid residues, which are simplified as two HTyr* and an Asp*, in a series of ligand exchange reactions. The calculations identified the Tyr* \rightarrow Tyr* \rightarrow Asp* (YYD) pathway as the optimal pathway. Along this pathway, the HTyr* residues may grab the free UO₂(CO₃)₃⁴⁻ aqueous species, and the proton transferred from HTyr* to the bound CO₃²⁻ ligand triggers the ligand exchange reactions. The Asp* residue builds interaction with uranyl in the end to anchor the uranyl in the binding site. This is also consistent with the experimental hypothesis that the Asp is relevant to the semi-open conformation of the uranylsTf complex.

The other two pathways, i.e. $Tyr^* \rightarrow Asp^* \rightarrow Tyr^*$ (YDY) and Asp* → Tyr* → Tyr* (DYY), in which the negatively charged Asp* coordinates to uranyl prior to the binding of one or both neutral HTyr* ligands, suggests that the excess ligands CO₃²⁻ to be replaced play a key role in determining the uptake mechanism of uranyl by sTf. The deprotonated Asp* does not have a proton to perturb the strong interaction between ${\rm CO_3}^{2-}$ and uranyl, which makes it energy demanding to repel one of the CO₃²⁻ ligands in order to release coordination site. In addition, in its carbonate complex, the positive charge of uranyl is already neutralized by the carbonate ligands, and the approaching of Asp* also brings electrostatic repulsive interaction, which disfavor its binding when uranyl is tri- or di-carbonated.

In summary, this work identified an optimal pathway for the uptake of uranyl by sTf at around pH = 7.0, and clarified that this is determined by synergistic ligands ${\rm CO_3}^{2-}$. This shows that at low pH, due to the strong preference of carbonate to be protonated, the UO₂(CO₃)₃⁴⁻ is no longer the predominant species,47 and the YYD mechanism may no longer dominate. This work thus helps improve our understanding on the *in vivo* migration of uranyl by providing detailed mechanism at the molecular level, and sheds light on the protein engineering and the design of decorporation reagents toward the enhanced capture of uranyl in specific pH range.

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References

- 1 J. Grenthe, J. Drożdżyński, T. Fujino, E. C. Buck, T. E. Albrecht-Schmitt and S. F. Wolf, in The Chemistry of The Actinide and Transactinide Elements, ed. L. R. Morss, N. M. Edelstein, J. Fuger and J. J. Katz, Springer, Dordrecht, The Netherlands, 2006.
- 2 M. Hada and A. G. Georgakilas, J. Radiat. Res., 2008, 49, 203-210.
- 3 Y. Knöbel, M. Glei, A. Weise, K. Osswald, A. Schäferhenrich, K. K. Richter, U. Claussen and B. L. Pool-Zobel, Toxicol. Sci., 2006, 93, 286-297.
- 4 S. S. Wise, W. D. Thompson, A. M. Aboueissa, M. D. Mason and J. P. Wise, Chem. Res. Toxicol., 2007, 20, 815-820.
- 5 P. Lestaevel, P. Houpert, C. Bussy, B. Dhieux, P. Gourmelon and F. Paquet, Toxicology, 2005, 212, 219-226.
- 6 P. Lestaevel, E. Romero, B. Dhieux, H. B. Soussan, H. Berradi, I. Dublineau, P. Voisin and P. Gourmelon, Toxicology, 2009, 258, 1-9.

Paper

7 O. Averseng, A. Hagege, F. Taran and C. Vidaud, Anal. Chem.,

2010, **82**, 9797–9802. 8 M. G. Benavides-Garcia and K. Balasubramanian, *Chem. Res.*

Toxicol., 2009, 22, 1613-1621.

- 9 M. Sutton and S. R. Burastero, *Chem. Res. Toxicol.*, 2004, 17, 1468–1480.
- 10 J. D. Van Horn and H. Huang, *Coord. Chem. Rev.*, 2006, **250**, 765–775.
- 11 G. Montavon, C. Apostolidis, F. Bruchertseifer, U. Repinc and A. Morgenstern, *J. Inorg. Biochem.*, 2009, **103**, 1609–1616.
- 12 S. Scapolan, E. Ansorborlo, C. Moulin and C. Madic, *Radiat. Prot. Dosim.*, 1998, **79**, 505–508.
- 13 M. Hémadi, N. T. Haduong, S. Plantevin, C. Vidaud and J. M. Chahine, *J. Biol. Inorg. Chem.*, 2010, 15, 497–504.
- 14 H. Sun, H. Li and P. J. Sadler, *Chem. Rev.*, 1999, **99**, 2817–2842.
- 15 M. Hémadi, N. T. Haduong and J. M. Chahine, J. Phys. Chem. B, 2011, 115, 4206–4215.
- 16 R. T. Mac-Gillivray, S. A. Moore, J. Chen, B. F. Anderson, H. Baker, Y. Luo, M. Bewley, C. A. Smith, M. E. Murphy and Y. Wang, *Biochemistry*, 1998, 37, 7919–7928.
- 17 H. M. Baker, Q. He, S. K. Briggs, A. B. Mason and E. N. Baker, *Biochemistry*, 2003, **42**, 7084–7089.
- 18 J. Wally, P. J. Halbrooks, C. Vonrhein, M. A. Rould, S. J. Everse, A. B. Mason and S. K. Buchanan, *J. Biol. Chem.*, 2006, 281, 24934–24944.
- 19 F. B. Abdallah and J. M. Chahine, *J. Mol. Biol.*, 2000, **303**, 255–266.
- 20 Z. Chikh, N. T. Haduong, G. Miquel and J. M. Chahine, J. Biol. Inorg. Chem., 2007, 12, 90–100.
- 21 Z. Chikh, M. Hémadi, G. Miquel, N. T. Haduong and J. M. Chahine, J. Mol. Biol., 2008, 380, 900–916.
- 22 N. T. Haduong, M. Hémadi, Z. Chikh and J. M. Chahine, *Biochem. Soc. Trans.*, 2008, 36, 1422–1426.
- 23 C. Vidaud, S. Gourion-Arsiquaud, F. Rollin-Genetet, C. Torne-Celer, S. Plantevin, O. Pible, C. Berthomieu and E. Quéméneur, *Biochemistry*, 2007, 46, 2215–2226.
- 24 M. G. Benavidesgarcia and K. Balasubramanian, *Chem. Res. Toxicol.*, 2009, **22**, 1613–1621.
- 25 O. Pible, P. Guilbaud, J. L. Pellequer, C. Vidaud and E. Quéméneur, *Biochimie*, 2006, **88**, 1631–1638.
- 26 M. Bühl, H. Kabrede, R. Diss and G. Wipff, *J. Am. Chem. Soc.*, 2006, **128**, 6357–6368.

- 27 S. O. Odoh, S. M. Walker, M. Meier, J. Stetefeld and G. Schreckenbach, *Inorg. Chem.*, 2011, 50, 3141–3152.
- 28 N. Kaltsoyannis, Chem. Soc. Rev., 2003, 32, 9-16.
- 29 D. Wang and Z. Chai, Chem. Soc. Rev., 2012, 41, 5836-5865.
- 30 W. Ding, W. Fang, Z. Chai and D. Wang, *J. Chem. Theory Comput.*, 2012, **8**, 3605–3617.
- 31 W. Ding, W. Fang, Z. Chai and D. Wang, *Acta Phys.-Chim. Sin.*, 2015, **31**, 1283–1301.
- 32 A. D. Becke, J. Chem. Phys., 1993, 98, 5648-5652.
- 33 S. Vosko, L. Wilk and M. Nusair, *Can. J. Phys.*, 1980, **58**, 1200–1211.
- 34 C. Lee, W. Yang and R. G. Parr, Phys. Rev. B: Condens. Matter, 1988, 37, 785–789.
- 35 B. Miehlich, A. Savin, H. Stoll and H. Preuss, *Chem. Phys. Lett.*, 1989, 157, 200–206.
- 36 M. J. Frisch, *et al.*, *Gaussian 09*, Gaussian Inc., Wallingford, CT, 2009.
- 37 W. Küchle, M. Dolg, H. Stoll and H. Preuss, *J. Chem. Phys.*, 1994, **100**, 7535–7542.
- 38 X. Cao, M. Dolg and H. Stoll, *J. Chem. Phys.*, 2003, **118**, 487–496.
- 39 X. Cao and M. Dolg, J. Mol. Struct., 2004, 673, 203-209.
- 40 K. Fukui, Acc. Chem. Res., 1981, 14, 363-368.
- 41 D.-C. Fang, Beijing Normal University, Beijing 100875, People's Republic of China, free of charge for academic users. Solution-phase free energy has been obtained from the THERMO program by using solution-phase translational entropy, which is different from that of the Gaussian 09 output based on gas-phase translational entropy.
- 42 M. Cossi, V. Barone, R. Cammi and J. Tomasi, *Chem. Phys. Lett.*, 1996, 255, 327–335.
- 43 G. Scalmani and M. J. Frisch, *J. Chem. Phys.*, 2010, **132**, 114110.
- 44 G. R. Grimsley, J. M. Scholtz and C. N. Pace, *Protein Sci.*, 2006, **15**, 1214–1218.
- 45 A. O. Tirler and T. S. Hofer, *J. Phys. Chem. B*, 2014, **118**, 12938–12951.
- 46 S. Tsushima, Y. Uchida and T. Reich, *Chem. Phys. Lett.*, 2002, 357, 73–77.
- 47 T. Lan, Z. Liu and D. Wang, Chin. J. Inorg. Chem., 2015, 31, 1774–1784.