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Role of EDTA protonation in chelation-based removal of mercury ions from water†

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A robust method of hazardous metal ion removal from an aqueous environment involves the use of chelating agents, such as ethylenediaminetetraacetic acid (EDTA). Here, we focus on mercury (Hg^{2+}) uptake by EDTA using both molecular dynamics and density functional theory simulations. Our results indicate that the deprotonation of the EDTA carboxylate groups improves the localization of negative charge on the deprotonated sites. This mechanism facilitates charge transfer between the metal ions and EDTA, and provides a stronger and more stable EDTA-Hg^{2+} complex formation improving the efficiency of the chelation process. The best metal removal conditions are achieved using the fully deprotonated form of EDTA, which naturally occurs at pH levels above 3.

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1 Introduction

Despite the rules imposed on mercury application, some industrial activities still cause mercury (Hg) pollution. Examples include paint and paper manufacturing, mining, coal burning, and the incineration of waste.^{1–3} Pollution can endanger the health of both humans and animals by poisoning vital organs.^{4–6} As a result, the World Health Organization and the environmental protection agency have listed mercury as one of the most toxic heavy metals.^{7–9}

The toxicity issue is further exacerbated by the fact that mercury can be found in different forms¹⁰ and sources, from the atmosphere to soil and groundwater.¹¹ This has motivated

the removal of mercury using different techniques,¹² for example coagulation and flocculation,^{13,14} precipitation,¹⁵ ion exchange,¹⁶ reverse osmosis,^{17,18} membrane filtration,^{19–21} and adsorption. Adsorption is the most attractive approach because of its simplicity and low cost.²²

Adsorption of mercury is typically based on chelation by a chelator (or a chelating agent) with two or more groups that are able to donate electron pairs for forming coordination bonds with metal ions and create a ring-like structure known as a chelate ring.²³ The most commonly used chelators for Hg removal and preconcentration are ethylenediaminetetraacetic acid (EDTA),^{24–26} hydroxyethylethylenediaminetriacetic acid (HEDTA),²⁷ 2,3-dimercaptopropanesulfonic acid (DMPS),²⁸ and 2,3-dimercaptosuccinic acid (DMSA).²⁹ In this work, we focus on EDTA. EDTA not only adsorbs many metal ions, such as mercury (Hg^{2+}), iron ($\text{Fe}^{2+}/\text{Fe}^{3+}$), and calcium ions (Ca^{2+}),^{30–34} but also outperforms many other commercial chelators in terms of adsorption energy/reactivity.³⁵

The understanding of Hg uptake by EDTA can help to improve the efficiency of chelation mechanisms in wastewater treatment. While some insight about such mechanisms is already available,^{35–37} the chelation mechanism is not fully understood.

To date, an EDTA-Hg^{2+} structure containing fully deprotonated EDTA, a Hg^{2+} ion and a single molecular water in the form of a distorted octahedral has been proposed using an X-ray absorption spectroscopy.³⁸ Furthermore, some density functional theory (DFT)-based studies have addressed the interaction of various metal ions with EDTA. The studies have, for example, focused on Na, K, Rb, Mg, Mn, Ca, and Sr,³⁹ Ba, Y and Zr,⁴⁰ Al, Sc, and V-Co,⁴¹ and Fe, Co, Cu.⁴² The interactions between Hg^{2+} and EDTA have also been studied³⁷ by applying

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† Electronic supplementary information (ESI) available: The force field parameters used in the molecular dynamics simulations are given in Tables S1–S5. The radial distribution functions and running coordination numbers obtained for the theoretical series of EDTA molecules are shown in Fig. S3–S7. The DFT results at different computational levels are presented in Table S6, along with the root-mean-square deviation of the structures from the experimentally resolved Hg^{2+} -EDTA-water complex. The corresponding structures are shown in Fig. S2, while the partial atomic charges of EDTA and Hg^{2+} -EDTA complexes at the different protonation states are shown in Fig. S8 and S9. The atom coordinates for the considered configurations are provided in the last section. See DOI: <https://doi.org/10.1039/d4cp02980a>

the B3LYP/LANL2DZ DFT level to some experimental structural data,^{43–45} suggesting the positive influence of K^+ presence on the complexation process. The impact of water on EDTA complex formation with some metals (other than Hg^{2+}) has also been addressed. For instance, some quantum mechanical/molecular mechanical (QM/MM) simulations⁴⁶ have revealed that Mg^{2+} binds to EDTA in a 6-fold octahedral complex, whereas Ca^{2+} prefers the formation of a 7-fold coordination complex with rapidly exchanging water molecules. In another work,⁴⁷ a combination of classical molecular dynamics simulations and extended X-ray absorption fine structure (EXAFS) has elucidated the impact of protonation on the binding of Eu^{3+} to EDTA in aqueous solutions. An additional example is DFT-based *ab initio* molecular dynamics simulation of EDTA complexation with Ln^{3+} , La^{3+} , Eu^{3+} and Gd^{3+} ions in aqueous solution at different pHs and temperatures.⁴⁸

Considering the possibility of water impact on the Hg^{2+} –EDTA complex formation and the necessity of understanding metal ion chelation mechanisms, in this study, we apply both DFT calculations and atomistic molecular dynamics (MD) simulations to investigate the process in aqueous environment. We consider both realistic and theoretical EDTA protonation scenarios to highlight the impact of both amino and carboxylate protonation on Hg^{2+} –EDTA complex formation. This approach not only guarantees the realistic study of the complex formation process over a wide pH range but also adds to the computational merit of the work by following a wide spectrum of changes in the protonation and charge distribution of EDTA. In the framework of our study, the MD simulations are particularly designed to sample all potential configurations of a complex and extract the most probable structures. The obtained Hg^{2+} –EDTA structures and the corresponding adsorption energies are then validated through DFT simulations to study the impact of charge distribution/protonation state on the efficiency of the chelation process.

2 Computational details

The uptake of Hg^{2+} by EDTA was studied using MD simulations and DFT calculations. The MD simulations were performed to

follow the uptake processes in explicit water solvent. The DFT simulations were used as a higher-level computational technique to re-evaluate the uptake behavior with higher accuracy. Using both approaches, we considered a number of protonation states for EDTA to study the impact of charge distribution on the uptake process. To include a wide range of protonation states in our studies, we considered two series of protonation states – one realistic and one theoretical – with the fully deprotonated EDTA^{4−} state as their starting point. For the theoretical states, we protonated one to four carboxylate groups, and distinguished them from the realistic protonation states by adding a 'th' prefix to their names. These structures are schematically shown in Fig. 1, where *th*EDTA⁰ is the neutral structure with 4 protonated carboxylate groups, and *th*EDTA^{1−}, *th*EDTA^{2−}, *th*EDTA^{3−} correspond to the different deprotonation levels with 3, 2, and 1 protonated carboxylate groups, respectively. We note that the *th*EDTA^{2−} can have two forms, which are labeled as *th*EDTA_a^{2−} and *th*EDTA_o^{2−}.

The inclusion of the *th*EDTA protonation states helps to isolate the impact of carboxylate protonation on the charge distribution and the complex formation process. However, such a protonation trend is only theoretical as EDTA protonation actually begins with the protonation of the amino groups rather than the carboxylates.⁴⁹ To address realistic conditions, we are concerned with the EDTA protonation states associated with various pH conditions. EDTA molecules in a complex with a heavy metal ion can exist in its fully deprotonated state at above pH 3.^{38,47,50,51} This is because, over pH 3, the proton of EDTA dissociates into the bulk solution, leaving the molecules in its fully deprotonated state (Hg^{2+} –EDTA^{4−}).^{38,47,52} Around pH 3–4, the EDTA protonation process starts, and Hg^{2+} –EDTA^{3−} complexes with one protonated amino group purple are also present in the solution, while the concentration of Hg^{2+} –EDTA^{4−} drastically decreases upon lowering pH.^{38,47} At low pH, the second amino protonation step takes place (Hg^{2+} –EDTA^{2−}),⁴⁷ resulting in a zwitterionic neutral state when complexed with divalent metal ions. Carboxylate protonation starts at very low pH conditions,⁴⁹ which is not relevant to industrial metal ion removal. Therefore, we added the EDTA^{3−} and zwitterionic EDTA^{2−} with one and two protonated amino groups, respectively, to our studied protonation states. Their corresponding schematic structures are shown in Fig. 1.

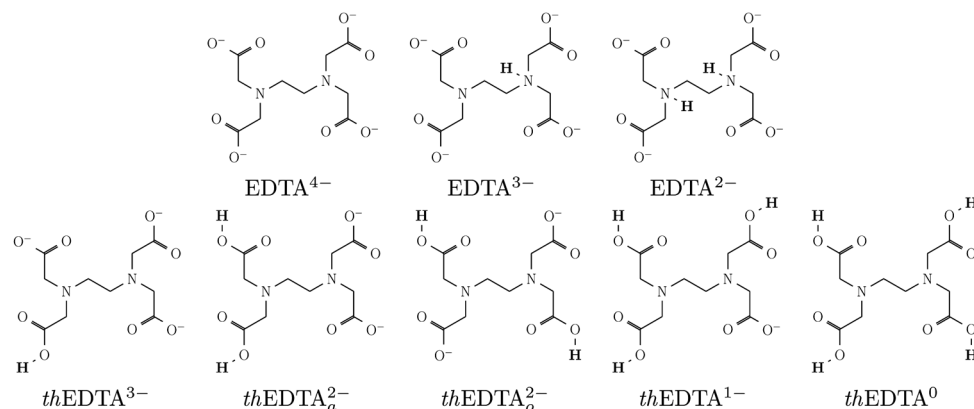


Fig. 1 Schematic view of the chemical structure of EDTA at the examined protonation states.

2.1 Molecular dynamics simulations

The formation of the aqueous mercury ion (Hg^{2+}) complexes with the EDTA molecules was investigated through molecular dynamics simulations using the LAMMPS software (version 2Aug2023).⁵³ To describe the interactions between the atoms, the OPLS/AA force field⁵⁴ and the SPC/E model⁵⁵ were applied to the EDTA and water molecules, respectively. For the mercury ion, the OPLS/AA and SPC/E-consistent parameters were taken from ref. 56–58. The adopted parameters are summarized in Tables S1–S5 (ESI†). The Moltemplate software⁵⁹ was used to generate the molecular structure of EDTA and the related input files.

The simulations were performed in four stages: (1) simulation of EDTA and Hg^{2+} chelation at constant volume and temperature $T = 298.15$ K (the NVT ensemble) to form a stable complex in the absence of water, while maintaining the total electroneutrality of the system by placing two counterions at fixed distances relative to the Hg–EDTA complex; (2) insertion of water molecules in the system obtained at the previous stage and running simulation at the constant pressure 1 atm and temperature $T = 298.15$ K (the NPT ensemble) until reaching the appropriate system density; (3) equilibration of the system obtained at the previous stage at the same NPT conditions; and (4) the production run at the same conditions and storing the trajectory for further analysis. During all these steps, EDTA and the counterions were kept apart at significant distances to focus exclusively on the EDTA–Hg complex and to avoid any unnecessary interaction between the complex and the counterions.

In stage 1, the system components were introduced in a cubic box of size $L_x = 150$ Å, $L_y = L_z = 75$ Å. EDTA was initially positioned at the maximum distance from Hg^{2+} . No water molecules were added to the box. To maintain system electroneutrality, two counterions were placed in the box at a distance larger than 60 Å from EDTA and Hg^{2+} . The distance was sufficient to minimize the effect of any counterion interactions with the complex. Undoubtedly, in the absence of water this effect remained significant. As such, all ions (*i.e.*, Hg^{2+} and the counterions) were immobilized during the simulation, but EDTA could move. The MD simulation was performed in the NVT ensemble at 298.15 K and resulted in the formation of a stable EDTA– Hg^{2+} complex within the time scale of 100 ps. The time step was 0.5 fs. The Nosé–Hoover thermostat was used to maintain the temperature using the relaxation parameter of 10 fs.

The system obtained from stage 1 was then immersed in 20 000 randomly positioned water molecules for stage 2. The minimum overlap distance between the molecules during the insertion was 1.5 Å. The aqueous system was then relaxed at $T = 298.15$ K and $P = 1$ atm using a relatively short simulation (100 ps) to achieve the expected density. The Nosé–Hoover thermostat-barostat with the relaxation parameters of 0.1 ps and 1 ps were respectively used to maintain the temperature and pressure. The position of Hg^{2+} was unfixed to allow the complex to move freely, but the position of the counterions remained fixed. At the end of this stage, the size of the simulation box decreased to $L_x = 134$ Å, $L_y = L_z = 67$ Å, and the density of water reached the value of 0.99 g cm^{−3}.

Because of the decrease in the simulation box size and the motion of the complex, the distance between the complex and the counterions decreased. However, it remained large enough (over 50 Å) to avoid any significant counterion–EDTA interaction.

In stage 3, the resultant system was equilibrated for the longer period of 1 ns to ensure stability at the ambient conditions. The other simulation details were identical to stage 2. The distance between the complex and the counterions was still large enough.

Finally, in stage 4, we performed 5 ns of the production run under the same conditions addressed in stages 2 and 3. At this stage, we monitored the positions of the complex and the counterions to ensure sufficient separation between them. According to our observations, the complex needed over 5 ns time to diffuse toward the counterions. Stage 4 generated a series of atomic position trajectories, which were stored every 1 ps for further analysis.

In all MD simulations, the periodic boundary conditions were applied in all directions. The long-range electrostatic interactions were computed using the particle–particle–particle–mesh (PPPM) scheme with 0.0001 accuracy and 12 Å cutoff radius. The cutoff of the Lennard-Jones (LJ) interactions was set to 10 Å. The bond lengths and angles of the rigid SPC/E water molecules were constrained using the SHAKE technique. The LJ cross-interactions were computed through the geometrical mixing rules. The non-bonding intramolecular interactions were considered for the 1–4 adjacent atoms of EDTA and scaled by a factor of 0.5 according to the OPLS/AA force field.

2.2 DFT calculations

The DFT calculations were performed using Gaussian 16 Rev. C1.⁶⁰ The stability of the structures was validated by the absence of any imaginary frequencies. To calibrate the required parameters, we initially considered several DFT approximations suggested for the simulation of ion uptake by EDTA and similar chelators,^{61–67} including B3LYP,^{68,69} M05-2X,⁷⁰ PBE,⁷¹ PBE0,⁷² and ω B97X-D.⁷³ HSE06⁷⁴ was also included because of its potential to provide accurate electronic structures.⁷⁵ For basis sets, we chose the ones that were accessible in the Gaussian package and contained all parameters needed to describe the Hg^{2+} –EDTA complex, *i.e.*, Def2-TZVP,^{76,77} Def2-TZVPP,^{76,77} and LanL2DZ.^{78,79} The SMD,⁸⁰ CPCM,⁸¹ and PCM⁸² models were used to describe the water environment implicitly.

The experimentally resolved Hg^{2+} –EDTA^{4−}–water structure³⁸ was optimized at different computational levels, and the resultant geometry was compared to the experimental counterpart. The comparison was quantified as root mean square displacement (RMSD) of the backbone structure by ChemCraft 1.8.⁸³ The results are summarized in Table S6, ESI† (along with the geometries displayed in Fig. S2, ESI†). The M05-2X/LanL2DZ/SMD scheme was found to be the most accurate computational level with the lowest RMSD value (0.694 Å). Consequently, we adopted this level for our DFT simulations. We noticed some discrepancies between DFT and the experiment regarding the attachment of the water to the complex. Therefore, we did not focus only on the experimental structure but considered the

possibility of alternative structures and the possible merits of the other DFT approximations. In the following, we started with the structures of the complex obtained from the MD simulations and refined them using different DFT computational levels.

3 Results and discussion

The chelation process was evaluated from two main perspectives: the structure of the complexes and the impact of the charge distribution/protonation state on their stability. The mechanisms of complex formation and the corresponding structures revealed by the MD simulations are discussed in Section 3.1. The charge distribution variations and the complex formation energies obtained from the DFT calculations are presented in Section 3.2.

3.1 Complex formation scenarios

The mechanism of EDTA-Hg²⁺ complex formation was studied using MD simulations. Different EDTA protonation states were considered by protonating the amino and carboxylate groups of EDTA. The EDTA protonation/charge states included *th*EDTA⁰, *th*EDTA¹⁻, *th*EDTA_a²⁻, *th*EDTA_o²⁻, *th*EDTA³⁻, EDTA³⁻, EDTA²⁻ and EDTA⁴⁻, where *th*EDTA⁰ is the EDTA structure with fully protonated carboxylate groups, EDTA³⁻ and EDTA²⁻ are the structures with protonated amino groups and EDTA⁴⁻ is the fully deprotonated counterpart. In our notation of the *th*EDTA²⁻ structures, the 'a' and 'o' indices indicate the protonation of the carboxylic acid groups positioned adjacent or opposite to the ethylene bridge of the EDTA molecule. The snapshots of the observed complexes are shown in Fig. 2. The corresponding radial distribution functions and running coordination numbers averaged over the collected MD trajectories are presented in Fig. 3–6 and Fig. S3–S7 (ESI†), where N and N_p refer to deprotonated and protonated nitrogen atoms, respectively, Oc denotes oxygen atoms of the carboxylate groups, and Ow corresponds to oxygen atoms of water molecules. The radial

distribution functions indicate preferable distances at which the atoms are located with respect to each other. The running coordination numbers allow us to estimate the number of atoms in the first coordination shell of the Hg²⁺ ion. The *th*EDTA⁰ case is not reported, because it did not form a complex with Hg²⁺ in our simulations.

According to Fig. 3, the formation of a complex from EDTA⁴⁻ and Hg²⁺ results in a single sharp peak in the radial distribution function of Hg–N around 2.1 Å, indicating the strong binding of Hg²⁺ to the two EDTA nitrogen atoms. Also, the radial distribution function of Hg–Oc indicates the coordination of Hg²⁺ with four Oc atoms at the distance of 2.0 Å. This octahedral structure is the most stable EDTA-Hg²⁺ complex throughout the simulation runs. The observed relative stability is an important finding because EDTA is naturally observed in its fully deprotonated state at pH levels greater than 3.^{38,47,50,51}

Moving from the fully deprotonated EDTA⁴⁻ state to the EDTA³⁻ structure with one protonated amino group (Fig. 4) we observe that Hg²⁺ detaches from one nitrogen atom, while it sustains its strong interaction with the second amino group. At the same time, it binds with three Oc atoms and dissociates from one carboxylate group. These changes in the coordination shell provide an opportunity for two water molecules to enter the first coordination shell of Hg²⁺ (Fig. 6, left panel). The protonation of the second amino group and the creation of EDTA²⁻ (Fig. 5) causes further disturbance in the adsorption of Hg²⁺ by hindering Hg²⁺ binding to the amino groups. The hindrance pushes Hg²⁺ from the distance of 2.1 Å observed for the fully deprotonated EDTA to the distances above 3.5 Å. This does not affect the coordination of Hg²⁺ with the carboxylate groups. However, it frees up a coordination site to accommodate an additional water molecule (Fig. 6, right panel). Therefore, we can conclude that the protonation of the amino groups weakens the tight binding of Hg²⁺ to EDTA and promotes its hydration.

A similar effect can be observed by following the theoretical protonation trend, which refers to the EDTA structures with

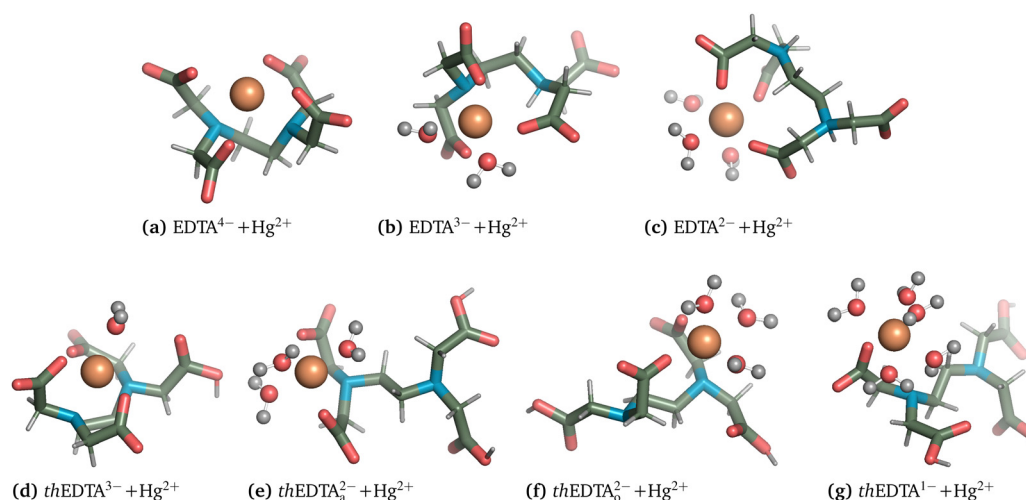


Fig. 2 The EDTA-Hg²⁺ complexes and the water molecules in their first coordination shell. The complexes are obtained from the MD simulations at ambient conditions. The green, blue, red, grey and orange colours represent the carbon, nitrogen, oxygen, hydrogen and Hg atoms, respectively.

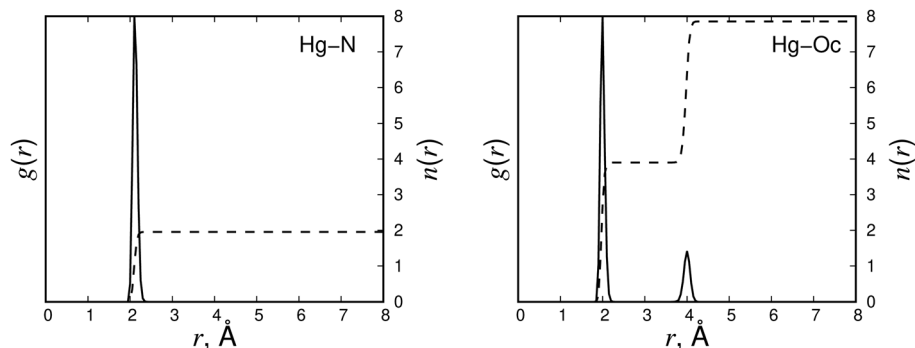


Fig. 3 The radial distribution functions $g(r)$ (solid line) and running coordination functions $n(r)$ (dashed line) for $\text{EDTA}^{4-} + \text{Hg}^{2+}$.

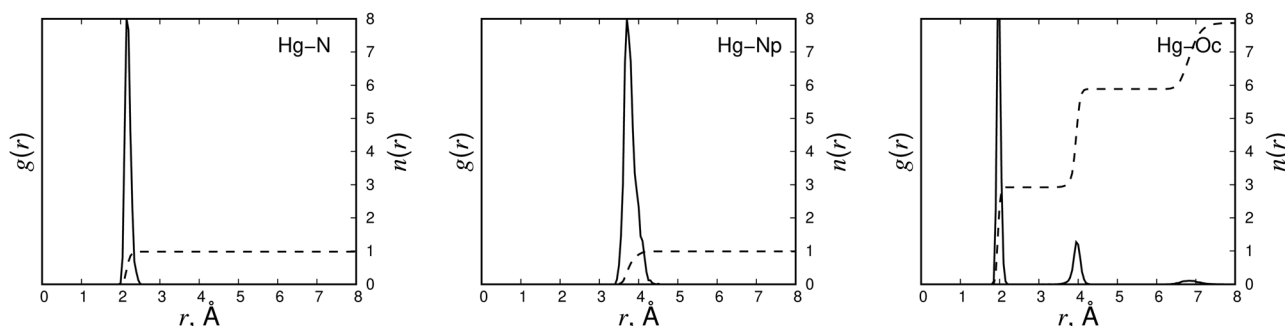


Fig. 4 The radial distribution functions $g(r)$ (solid line) and running coordination functions $n(r)$ (dashed line) for $\text{EDTA}^{3-} + \text{Hg}^{2+}$.

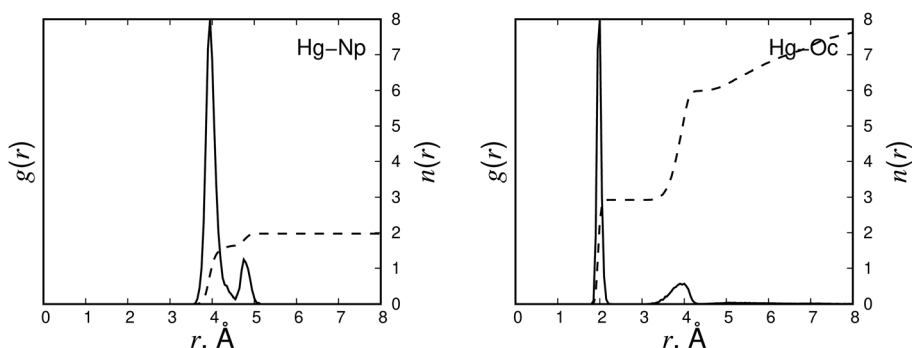


Fig. 5 The radial distribution functions $g(r)$ (solid line) and running coordination functions $n(r)$ (dashed line) for $\text{EDTA}^{2-} + \text{Hg}^{2+}$.

protonated carboxylate and deprotonated amino groups. According to Fig. S3 (ESI[†]), the protonation of one carboxylate group (thEDTA^{3-}), leads to the permanent positioning of Hg^{2+} at 2.0–2.5 Å from the two nitrogen atoms of EDTA, and about 2.0 Å from the three deprotonated oxygens.

When two adjacent carboxylate groups are protonated (thEDTA_a^{2-}), Hg^{2+} binds to the two deprotonated Oc atoms with an average bond length of 2 Å (Fig. S4, ESI[†]) and to one deprotonated nitrogen atoms at about 2.2 Å. The same is obtained by protonating two opposite carboxylate groups of EDTA ($\text{thEDTA}_o^{2-} + \text{Hg}^{2+}$), but the different conformation of EDTA molecule is observed in this case (Fig. S5, ESI[†]). It should be noted that we also found several alternative $\text{thEDTA}_o^{2-} - \text{Hg}^{2+}$

configurations, where an additional bond appeared between Hg^{2+} and an oxygen of one carboxylate group, as well as a bond between Hg^{2+} and the second nitrogen atom. Nevertheless, these configurations of the complex appeared to be short-lived, lasting less than 1.5 ns. The most frequently observed $\text{thEDTA}_o^{2-} - \text{Hg}^{2+}$ structure is illustrated in Fig. 2.

When three carboxylate groups are protonated (thEDTA^{1-}), the Hg^{2+} ion binds to the deprotonated carboxylate oxygen at around 2 Å (Fig. S6, ESI[†]). Hg^{2+} stays rather far from the nearest nitrogen atom at the distance of about 3.9 Å, as in the case of EDTA^{2-} with protonated amino groups.

As noted for the EDTA structures with protonated amino groups, the unbinding of Hg^{2+} from the N and Oc atoms in the

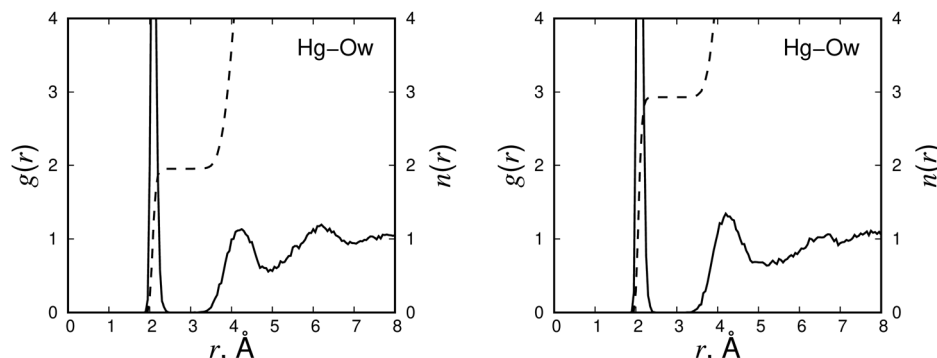


Fig. 6 The radial distribution functions $g(r)$ (solid line) and running coordination functions $n(r)$ (dashed line) of Hg^{2+} –Ow in the complexes of $\text{EDTA}^{3-} + \text{Hg}^{2+}$ (left panel) and $\text{EDTA}^{2-} + \text{Hg}^{2+}$ (right panel).

theoretical protonation series, enhances Hg^{2+} hydration. Based on Fig. S7 (ESI†) there are 1, 3, 1–3 (with 3 being the most probable), and 5 water molecules in the first coordination shell of Hg^{2+} in the thEDTA^{3-} , thEDTA_a^{2-} , thEDTA_o^{2-} and thEDTA^{1-} complexes, respectively. In the EDTA^{4-} – Hg^{2+} complex, the coordination shell of the ion contains no water molecules, in agreement with the DFT results presented in Fig. S2 (ESI†), displaying water detachment from Hg^{2+} after structural optimization using several DFT approximations.

The observed ion unbinding from the protonated N and Oc atoms and the increased ion hydration are in perfect agreement with the findings of Licup *et al.*⁴⁷ regarding Eu^{3+} chelation to EDTA. Using extended X-ray absorption fine structure (EXAFS) measurements at pH 3 and 11, Licup *et al.* observed the increase of Eu–N bond distances resulting from carboxylate protonation and the exposure of Eu^{3+} to water. They also reported full detachment of Eu^{3+} from fully protonated EDTA, which is consistent with the fact that a stable Hg^{2+} – thEDTA^0 complex was not observed in our MD simulations.

3.2 Complex formation energies

One can expect that the changes in the hydration shells and the structure of the complex implicate varying Hg^{2+} adsorption energies and, therefore, uptake efficiencies. To confirm this, we first calculated the formation energies of the complexes obtained from the MD simulations. Then, we exported the complexes to DFT simulations and reoptimized the complex structures (and their isolated constituting components) at different computational levels. The complexation Gibbs free energies (G_{cmplx}) were calculated as the difference between the energy of the complex ($G_{\text{EDTA}+\text{Hg}[+n\text{H}_2\text{O}]}$) and its isolated components including the Hg^{2+} ion (G_{Hg}), water ($G_{\text{H}_2\text{O}}$), and the corresponding the EDTA conformer (G_{EDTA}) through formula $G_{\text{cmplx}} = G_{\text{EDTA}+\text{Hg}[+n\text{H}_2\text{O}]} - G_{\text{EDTA}} - nG_{\text{H}_2\text{O}} - G_{\text{Hg}}$, where n is the number of explicit water molecules (see Fig. 2). The results are summarized in Table 1.

The MD results in Table 1 show that the higher the degree of EDTA protonation, the higher the Gibbs free energy of Hg^{2+} complex formation and the lower the stability of the complex. The results strongly suggest that the naturally occurring fully deprotonated EDTA structure provides the highest efficiency of

Hg^{2+} uptake by enabling an octahedral chelation mode (see Fig. 2). Similarly, the DFT results show a significant increase in the free energy of complex formation upon protonation of the amino and carboxylate groups of EDTA. All levels of theory present a similar trend, though they display some numerical differences resulted from various DFT approximations. They all suggest the fully deprotonated EDTA^{4-} state for Hg^{2+} removal. This suggestion is in line with the findings of earlier studies, which reported the formation of stable metal ion– EDTA^{4-} complexes for Ln,⁴⁸ and La, Ce, Pr, and Nd.⁸⁴ Furthermore, all results point out that even partial EDTA deprotonation facilitates Hg^{2+} uptake, guaranteeing Hg^{2+} chelation over a wide range of pH.

A question that needs addressing is how the protonation state of EDTA affects the energies. The adsorption energy calculated from MD simulations equals the difference between the potential energy of the Hg^{2+} ion in the complex and its potential energy in bulk water. In MD, these energies consist of van der Waals interactions (computed using LJ parameters) and electrostatic contributions (estimated using partial atomic charges). The OPLS/AA force field suggests interaction parameters that differ depending on the protonation/deprotonation of the carboxylate or amino groups of EDTA, as detailed in Table S1 (ESI†). Specifically, there are significant changes in the partial charges of the oxygen and nitrogen atoms belonging to these groups (*cf.* O1/O2 vs. O3, N1 vs. N2), which play a crucial role in Hg–EDTA complexation. However, in classical MD, these changes are not influenced by the conformation of EDTA or the arrangement of the Hg^{2+} ion and water molecules. Further insights into the charge distribution in the EDTA molecule

Table 1 G_{cmplx} (kJ mol^{−1}) calculated at different levels of theory

Complex	MD	B3LYP/ LanL2DZ	M05-2X/ LanL2DZ	ωB97X-D/ Def2-TZVPP
$\text{EDTA}^{4-} + \text{Hg}^{2+}$	−181.7 ± 33.2	−197.2	−223.6	−247.5
$\text{EDTA}^{3-} + \text{Hg}^{2+}$	−112.1 ± 33.7	−72.8	−100.4	−108.9
$\text{EDTA}^{2-} + \text{Hg}^{2+}$	−141.9 ± 35.8	−45.3	−92.6	−70.5
$\text{thEDTA}^{3-} + \text{Hg}^{2+}$	−115.4 ± 32.8	−143.5	−185.6	−191.1
$\text{thEDTA}_a^{2-} + \text{Hg}^{2+}$	−82.4 ± 33.1	−81.0	−82.5	−89.6
$\text{thEDTA}_o^{2-} + \text{Hg}^{2+}$	−77.6 ± 35.5	−60.3	−61.6	−88.3
$\text{thEDTA}^{1-} + \text{Hg}^{2+}$	−63.4 ± 35.3	−10.0	−18.6	26.6

can be provided by DFT calculations, which we performed for a series of protonation states of EDTA at the M05-2X/LanL2DZ level of theory.

According to the atomic polar tensor (APT) charges reported in Fig. S8 and S9 (ESI[†]), protonation changes the charge distribution throughout the EDTA molecule. The extent of the changes depends on the proximity of atoms to the protonated centres and the conformation of the EDTA molecule. In the deprotonated carboxylate groups, oxygen atoms have a larger negative charge compared with protonated ones. The absence of positively charged hydrogen atoms and more negatively charged oxygens of deprotonated carboxylate lead to strong binding of EDTA with the positive Hg^{2+} ion. The deprotonated nitrogen atoms in amino groups also bear significant negative charges, thus they are involved in binding with the ion as well. In a completely deprotonated EDTA molecule (EDTA^{4-}), this results in the formation of 6-fold coordination geometry around Hg^{2+} ion consisting of four carboxylate oxygen atoms and two amino nitrogen atoms, similar to that observed in MD simulations.

When one of the amino groups is protonated (EDTA^{3-}), the negative charge accumulated on the nitrogen atom decreases, and the attached positively charged hydrogen atom shields the amino group from electrostatic interaction with the Hg^{2+} ion. This obviously weakens the attractive interaction between the ion and the protonated nitrogen atom, causing dissociation between them. The same is observed in MD simulations, where the reduction in the charge of protonated nitrogen is even more pronounced compared to the corresponding DFT predictions. On the other hand, the DFT calculations show that interaction between EDTA molecule and Hg^{2+} ion induces the transfer of some additional negative charge to the negatively charged carboxylate and deprotonated amino atoms, which balances the excessive positive charge of the Hg^{2+} ion acquired through its interaction with certain oxygen atoms of the water molecules. Simultaneously, the carboxylate oxygen atoms of EDTA and the hydrogen atoms of the water molecules can form hydrogen bonds and transfer some charge to stabilize the generated EDTA–Hg–water complex. It is observed, that only three oxygen atoms of carboxylate groups remain in the first coordination shell of the ion, what is in agreement with our MD simulations.

The ion dissociates from both nitrogen atoms when two amino groups are protonated (EDTA^{2-} – Hg^{2+}). Due to this, the structure of the Hg–EDTA complex becomes distorted, and Hg^{2+} ion binds only with one carboxylate group of EDTA. Two other bonds are created between the ion and the oxygen atoms of water molecules, which, at the same time, are bound to carboxylate groups through hydrogen bonds. Contrary to this, in MD simulations, the Hg^{2+} ion binds directly to three carboxylate groups.

Similarly, in the theoretical series, we observe significant changes in the partial charge of the protonated oxygen atoms. The changes are more significant for the protonated oxygen atom and lead to the localization of less negative charge on them. This is simultaneous with a slight decrease in the charge

of the adjacent carbon atom, as well as a slight increase in the charge of the closest nitrogen atom. These changes weaken the electrostatic interaction of Hg^{2+} with the negatively charged centres, transferring less charge during the process and reducing the stability of the resultant complex. The charge transfer process manifests itself by increasing the partial charge of Hg^{2+} and the carbon atom bonded to the deprotonated oxygen atom and reducing the partial charge of the deprotonated oxygen and nitrogen atoms.

Therefore, the charge analysis of both theoretical and realistic EDTA protonation series ascertains that a higher degree of deprotonation can guarantee stronger binding of the mercury ion with the chelating agent. The deprotonated carboxylate groups and the deprotonated amino groups atoms are preferred for binding between the Hg^{2+} ion and EDTA molecule, offering more negatively charged centres for electrostatic interaction.

4 Conclusions

The combination of the MD and DFT simulation approaches demonstrates the importance of EDTA deprotonation in the uptake of mercury ions from water. According to our results, the availability of more deprotonated amino and carboxylate sites facilitates mercury interaction with EDTA, leading to stronger adsorption and a higher efficiency of mercury removal. As EDTA molecule complexed with mercury ion predominantly exists in the fully deprotonated state over a wide range of pH (greater than 3), its high chelation efficiency is expected at both alkaline and acidic conditions. Our findings about the effect of EDTA protonation on Hg–EDTA complex formation agree with earlier studies on EDTA chelation to other metal ions, which also highlight protonation effects on similar chelating agents such as HEDTA, NTA, DTPA, HEIDA, and GLDA.⁸⁵ The control of pH conditions, and thus the protonation and charge distribution of chelating agents, is a crucial factor in optimizing chelation processes. The most efficient protonation state can be determined through both simulation and experimental methods.

Data availability

The data supporting this article have been included as part of the ESI.[†]

Conflicts of interest

There are no conflicts to declare.

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