Interplay of Physically Different Properties Leading to Challenges in Separating Lanthanide Cations – an Ab Initio Molecular Dynamics and Experimental Study

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Abstract

The lanthanide elements have well-documented similarities in their chemical behavior, which makes the valuable trivalent lanthanide cations (Ln$^{3+}$) particularly difficult to separate from each other in water. In this work, we apply ab initio molecular dynamics simulations to compare the free energies ($\Delta G_{\text{ads}}$) associated with the adsorption of lanthanide cations to silica surfaces at a pH condition where SiO$^-$ groups are present. The predicted $\Delta G_{\text{ads}}$ for lutetium (Lu$^{3+}$) and europium (Eu$^{3+}$) are similar within statistical uncertainties; this is in qualitative agreement with our batch adsorption measurements on silica. This finding is remarkable because the two cations exhibit hydration free energies ($\Delta G_{\text{hyd}}$) that differ by $>2$ eV, different hydration numbers, and different hydrolysis behavior far from silica surfaces. We observe that the similarity in Lu$^{3+}$ and Eu$^{3+}$ $\Delta G_{\text{ads}}$ is the result of a delicate cancellation between the difference in Eu$^{3+}$ and Lu$^{3+}$ hydration ($\Delta G_{\text{hyd}}$), and their difference in binding energies to silica. We propose that disrupting this cancellation at the two end points, either for adsorbed or completely desorbed lanthanides (e.g., via nanoconfinment or mixed solvents), will lead to effective Lu$^{3+}$ separation.
I. INTRODUCTION

Lanthanide (Ln) series elements exhibit specific utility in green energy applications including lighting, wind turbines, electrified vehicles, and catalysis. Improved mining and extraction techniques are needed to expand the inventory of these critical elements. Trivalent lanthanide cations (Ln$^{3+}$) naturally occur as mixtures. As such, separation of Ln$^{3+}$ from each other is a technologically relevant and chemically challenging problem. Ln separation schemes are complicated and hazardous, and have to be adjusted depending on the composition of the Ln-containing ore. Ion-exchange separation of lanthanides has limited industrial use due to separation efficiencies of existing resins not yielding sufficient Ln$^{3+}$ separation. Here we investigate what makes adsorption more or less selective, so in the future we can develop ion-exchange separation methods based on the fundamental science insight.

This separation problem has been addressed from several angles. Extensive basic science research has been conducted to elucidate the solvation properties of lanthanides in liquid water. Further experimental and theoretical work has focused on separating Ln$^{3+}$ from their mixtures using organic ligands. The results suggest that factors like pH, counter-ion effects, soft-hard ion concepts, and ligand rigidity are important. Additional studies have focused on the possibility of using silica and nanoporous silica materials (with and without functionalization of their surfaces) to adsorb and separate Ln$^{3+}$. This latter pathway has proven somewhat successful. At pH $\sim$4, bare silica nanopores have been shown to selectively adsorb scandium, but selectivity among lanthanides appears limited. Ilgen has shown that, at pH $\sim$6, smaller lanthanides cations are preferentially retained on nanoporous silica, depending on the nanopore size. At still higher pH values, lanthanides are less soluble and precipitate as hydroxides. Improvement of such inorganic materials will likely enhance their Ln$^{3+}$ selectivity, making them competitive with organic materials which are more costly, less durable, and less environmentally benign.

Further studies are needed to determine how separation mechanisms determined for organic ligands can be transferred to separation using inorganic materials. Computationally, molecular-level mechanistic studies have had difficulty determining lanthanide bonding to inorganic surfaces or the relevant adsorption/desorption free energies ($\Delta G_{ads}$), in part because of a lack of classical force fields that accurately reflect the interaction between $f$-electron trivalent cations and water or silica surfaces.

In this work, we address the fundamental question of why different Ln$^{3+}$ cations exhibit “chemically similar” adsorption equilibrium constants on silica within a specific pH range, despite their significantly different physical properties. For example, lanthanides are known to exhibit hydration free energies in liquid water ($\Delta G_{hyd}$) that differ significantly, by $>$2 eV. Lutetium (Lu$^{3+}$) and europium (Eu$^{3+}$) are at the endpoint and midpoint of the $4f$-electron block of the periodic table, respectively. Their ionic radii, differing by $\sim$0.1 Å, are suf-
ficient to cause the Lu$^{3+}$ $\Delta G_{\text{hyd}}$ to be more negative (i.e., favorable) than that of Eu$^{3+}$ by $\Delta \Delta G_{\text{hyd}} = -2.17$ eV. We will show that this difference far exceeds the relative adsorption free energy ($\Delta \Delta G_{\text{ads}}$). We will also demonstrate that the two Ln$^{3+}$ exhibit different extent of hydrolysis reactions involving splitting of H$_2$O molecules in their inner spheres; such concerted hydrolysis/desorption behavior has been proposed and reported for other multivalent cations. As a result, Ln$^{3+}$ hydration numbers ($N_{\text{hyd}}$, the numbers of H$_2$O molecules coordinated to different Ln$^{3+}$) can differ by two along parts of the desorption reaction pathway.

We apply the ab initio molecular dynamics (AIMD) method, based on Density Functional Theory (DFT), in conjunction with the potential-of-mean-force (PMF) method which effectively extends AIMD time scales to yield accurate free energies. We show that Eu$^{3+}$ and Lu$^{3+}$ exhibit very similar $\Delta G_{\text{ads}}$. These predictions are supported by batch adsorption measurements on non-porous silica surfaces. The advantage of DFT is that it can be further applied to analyze different energy contributions. Our analysis leads us to conclude that Ln$^{3+}$ are not very similar; instead, the cancellation of large energy terms leads to similar $\Delta G_{\text{ads}}$ that hinders separation. From this work, we propose that the key to successful Ln$^{3+}$ separation technology resides in the disruption of this cancellation in either the adsorbed or desorbed regime.

This work involves significant computational challenges. The accuracy of DFT functionals and pseudopotentials used to treat Ln$^{3+}$ needs to be addressed, and we will perform tests on one of the Ln$^{3+}$ pseudopotential used in this work. However, there is an urgent need to address the accuracy of other modeling details as well, such the explicit treatment of both outer-shell solvating water molecules and the environment near interfaces. In implicit outer-shell solvation DFT calculations, the predicted $\Delta G_{\text{hyd}}$ exhibit variations that are on the order of a fraction of an eV. The discrepancies partly arise from the use of different DFT functionals or quantum chemistry methods, but variations in the implicit solvation methods used also likely play a role. In this work, we apply exclusively explicit hydration treatment via AIMD. We argue that such statistical mechanically rigorous research needs to take place in parallel with the development of more accurate DFT methods when dealing with Ln$^{3+}$ in aqueous interfacial environments. AIMD calculations of $\Delta G_{\text{ads}}$ at water/mineral interfaces for trivalent lanthanide cations, or for any trivalent cations, remain rare and challenging because of the high charge density involved. Our work represents a key step in this direction.

II. METHOD

A. Experimental Details

Adsorption of Eu$^{3+}$ and Lu$^{3+}$ onto amorphous silica was quantified. We used commercially available fumed silica (Sigma Aldrich) with surface area of 192±3 m$^2$ g$^{-1}$ as reported
in our earlier work. No background electrolyte or buffer were used. Milli-Q water with the resistivity of 18 MΩ·cm was used for all stock solutions and experiments. Lanthanide (Ln) stock solutions were prepared by diluting their nitrate salts Ln(NO$_3$)$_3$ in milli-Q water. Aqueous concentration was verified by inductively coupled plasma mass spectrometry (ICP-MS) analysis as described below.

Initially 50±1 mg of silica was weighted into each centrifuge vial, then 10 mL of Milli-Q H$_2$O was added, and samples were hydrated for a minimum of 2 hours. To begin the adsorption experiment, lanthanide stock solution was added, and the total volume of each sample was brought to 50 mL. The pH was measured after 2 hours of reaction and read at pH=5.0 ±0.3 for all samples. Then the samples were left on a shaker table for one-week (168 hours). Our preliminary kinetics runs showed that adsorption equilibrium is reached after 48 hours. The initial concentration of each lanthanide in the reactors was 0.1, 1, 10, 20, 30, and 50 µM L$^{-1}$. The sample with 10 µM L$^{-1}$ was made in triplicate to assess experimental error. All experiments were performed at ambient temperature (22 °C).

B. AIMD Computational Details

Finite temperature AIMD simulations apply the Perdew-Burke-Ernzerhof (PBE) functional, the projector-augmented wave-based Vienna Atomic Simulation Package (VASP), a 400 eV energy cutoff, and Γ-point sampling of the Brillouin zone. A Nose thermostat maintains the temperature at a slightly elevated 400 K. A Born-Oppenheimer energy convergence criterion of 10$^{-6}$ eV and a time step of 0.5 fs are enforced. These settings are similar to those in our previous ion desorption work. The charge-neutral simulation cell has a Si$_{40}$O$_{88}$H$_{3}^{+}$ stoichiometry for the reconstructed β-cristobalite (001) slab, 123 H$_2$O molecules, and a Lu$^{3+}$ or Eu$^{3+}$ cation in an initially bidentate adsorbed configuration. All simulation cells have dimensions 14.32Å×14.32Å×26.0 Å. They represent a 2$^{1/2}$×2$^{1/2}$ expansion of simulation cells we previously applied. The larger simulation cells are adopted because of the expectation that trivalent cations will experience stronger image-image interactions at the same cell size. The VASP lanthanide pseudopotential used are “Lu 23Dec2003” and “Eu 3 20Oct2008.” Lu$^{3+}$ has no unpaired f-electrons and the Eu pseudopotential adopted (henceforth referred to as “Eu(A)”) subsumes its f electrons into the core; hence non-spin polarized DFT is applied for all AIMD simulations. Calculations using the Lu pseudopotential, with explicit 4f electrons, are expected to be more accurate than those using the Eu(A) pseudopotential, without explicit 4f electrons. Some static, spin-polarized DFT, DFT+U, and HSE06 calculations, using the “Eu 23Dec2003” (“Eu(B)”) pseudopotential with an explicit, partially filled f-shell, are conducted as spot checks (Sec. IID).

The number of H$_2$O molecules in the simulation cell is determined as follows. Classical
force field-based grand canonical Monte Carlo (GCMC) simulations are first applied to
determine the average number of water molecules filling the gap between the silica surfaces.
The SPC/E water model, a force field for silica based on OPLS and generic force field
parameters pertinent to Ln$^{3+}$ are adopted for this purpose. Silica atoms and the adsorbed
cation are frozen in DFT-optimized positions in GCMC calculations; only water molecules
are inserted into or removed from the simulation cell. GCMC yields 7 H$_2$O molecules
coordinated to the Ln$^{3+}$ adsorbed to the surface. Switching to AIMD simulations and a
Lu$^{3+}$ cation reduces this to four after equilibration (Fig. S2a).

The AIMD calculations in this work omit dispersion corrections. This enables compar-
ison with our previous pK$_a$ predictions which involve a similar computational protocol.
Adding dispersion is known to improve AIMD predictions of liquid water structure ($g(r)$) at
T=300 K. But it has yet to be demonstrated that this gives universally superior predictions
at water/oxide interfaces.

In the presence of acid functional groups at water/material interfaces, the pH in the
simulation cell should be pinned at the pK$_a$ of functional groups, provided that (1) there is
only one type of such groups; (2) a fraction of them are deprotonated; (3) their pK$_a$ is lower
than that of H$_2$O; and (4) the surface groups do not interact with each other. Within the
non-interacting assumption, the pH in our AIMD cells should be between 7.0 and 8.1 – the
pK$_a$ range previously predicted for this single type of SiOH on this surface. In experimental
samples with amorphous or crystalline silica, bimodal or trimodal pK$_a$ distributions have
been reported. It would have been more challenging to assign pK$_a$ in AIMD simulation
cells with multiple types of SiOH.

C. Potential-of-Mean-Force Details, Reaction Coordinates

The potential-of-mean-force profile is computed as $\Delta W(Z) = -k_B T \log P(Z)$ where $P(Z)$
is the probability that a $Z$ value is recorded in the trajectory within a window, after mak-
ing adjustments to rigorously remove the effect of umbrella sampling penalties. Here $Z$ is
the coordinate normal to the silica-water interface, $Z=z_{Ln} - z_{Si}$, Ln is the desorbing lan-
thanide cation, and Si is the Si atom close to the two O$_{Si}^{-}$ groups initially coordinated to the
Ln$^{3+}$. This coordinate is chosen to accommodate the strong electrostatic attraction between
trivalent cations and surface silanol groups which can exhibit substantial bending motion. Harmonic penalties $A_o(Z - Z_i)^2$ are added to DFT energies in a series of windows with
a progression of $Z_i$ values, separated by 0.3 Å spanning the reaction paths. $A_o$ is set at
2 eV/Å$^2$.

For Lu$^{3+}$, the initial configuration in the window has the cation coordinated to two SiO$^-$
groups. Then AIMD is applied. Each subsequent window, with successively larger $Z_i$, and
therefore greater extent of desorption, is initiated by taking a configuration near the end
of the trajectory from the previous window along the Z-coordinate. The first one or more picoseconds in each window is used for equilibration only; statistics are collected for up to 45 ps. Statistical uncertainties in ΔW(Z) are estimated by splitting the trajectory in each window into five, calculating the standard deviation in ΔW(Z) between the edge Z values in each window (ΔΔW(Zi)), and propagating the noise across windows assuming gaussian statistics. For Eu$^{3+}$ simulations, each sampling window is initiated using an equilibrated configuration taken from the Lu$^{3+}$ trajectory with the same Zi value. This generally entails an increase in hydration number ($N_{\text{hyd}}$) by one compared to Lu$^{3+}$, and sometimes reduces the number of OH$^-$ groups coordinated to the Eu$^{3+}$ ($N_{\text{oh}}$). It typically takes 4-10 ps to equilibrate $N_{\text{hyd}}$ for Eu$^{3+}$ in each window. Trajectory lengths in different umbrella sampling window are listed in Table S1 in the ESI. The aggregate trajectory lengths used in all windows exceed 336 and 255 ps for Lu$^{3+}$ and Eu$^{3+}$, respectively.

In the $Z_i=5.0$ Å sampling window, the second of two O$_{\text{Si}}$-Ln$^{3+}$ ionic bond is being broken, forming two valleys in the free energy landscape with the absolute distance (not just z-coordinate) $R'=R_{\text{Ln-O}}$ centered around $R'=2.5$ Å (with one O$_{\text{Si}}$-Ln$^{3+}$ bond) and 4.2 Å (with zero, Sec. III B). These valleys are separated by a small free energy barrier. In sampling windows with $Z_i \ll 5.0$ Å, the $R' \sim 2.5$ Å valley is strongly favored, while large $Z_i$ strongly favors the $R' \sim 4.2$Å valley. As our reaction coordinate Z only constrains the vertical distance between Ln$^{3+}$ and the designated Si atom, it does not yield a smooth transition between the two $R'$ valleys in the handshake region near $R \sim 5.0$ Å (Sec. III B).

To deal with this problem and generate a smooth ΔW(Z), a secondary umbrella sampling calculation is performed on reaction coordinate $R'=R_{\text{Ln-O}}$, as follows. (a) By trial and error, we locate the sampling window (or create a new sampling window) centered around $Z=Z_i$ where the $R_{\text{Ln-O}} \sim 2.5$ Å and $R_{\text{Ln-O}} \sim 4.0$ Å valleys are similar in free energy. This occurs at $Z_i=5.0$ Å and 4.9 Å for Eu and Lu, respectively. (b) Keeping the primary umbrella sampling $A_o$ and $Z_i$ parameters constnat, we introduce a series of harmonic constraints $C_o (R'-R_i)^2$, with $C_o$ chosen to be 1.5 or 2.0 eV/Å$^2$ and $R_i$ separated by between 0.2 to 0.4 Å. (c) Ideally, one would generate a 2-dimensional PMF plot with $Z$ and $R'$. In reality, the relatively short AIMD simulations do not permit compiling accurate 2-D PMF statistics. Instead, we align the $R'$ windows by integrating all Z contributions in a restricted range that feature in the two end-point $R'$ sampling windows, so that there is overlapping statistics. The ranges chosen are $4.73$ Å < $Z$ < $4.83$ Å for Lu$^{3+}$ and $4.87$ Å < $Z$ < $4.97$ Å for Eu$^{3+}$. (d) Integrating the ΔW(Z, $R'$) over this narrow Z range generates a pre-factor $F$ that describes the statistical weight of the two $R'$ valleys. (d) If $F$ is smaller than $1/15$, we swtich to larger $Z_i$ and retry (a)-(d). If $F$ is larger than 15, we decrease $Z_i$ instead. As mentioned above, $Z_i=5.00$ Å and 4.90 Å are chosen for the Eu$^{3+}$ and Lu$^{3+}$ simulations via trial-and-error. (e) We add the probabilities $P(Z)$ from the two valleys, computed with $C_o=0$ (unconstrained in the $R'$ coordinate), weighted by the factor $F$.

A less severe version of this problem was encountered in our previous calculations asso-
associated with Cu\textsuperscript{2+} desorption.\textsuperscript{25} The Cu\textsuperscript{2+}-O(SiO\textsuperscript{−}) attraction is considerably weaker than the Ln\textsuperscript{3+}-O(SiO\textsuperscript{−}) attraction, and a less elaborate procedure was devised to circumvent this issue.\textsuperscript{25} The higher local charges associated with Ln\textsuperscript{3+} cations makes AIMD PMF desorption calculations more challenging.

Unlike Ref.\textsuperscript{25}, we do not use \( Z = (z_O - z_M) \) for reaction coordinate, where O is one of the O\textsubscript{SiO} groups initially coordinated to the metal (“M”) cation. The attraction between Ln\textsuperscript{3+} and the O atom in a surface SiO\textsuperscript{−} group is stronger than in previous divalent cation calculations. As a result, using the previous \( Z = (z_O - z_M) \) coordinate can lead to the O atom being pushed into the silica interior while the Ln\textsuperscript{3+} cation remains on the surface, bonded to other surface silanol groups. Unlike Ref.\textsuperscript{48}, we do not use the distance \( R = |R_O - R_M| \). This coordinate also allows the cation to roll along the surface on to different surface sites, instead of away from the surface into the bulk liquid. The water oxygen-cation coordination number \( N_{hyd} \) is not used for our purpose; it does not distinguish possible outer-sphere (Ln\textsuperscript{3+}/H\textsubscript{2}O/SiO\textsuperscript{−}) complexes, where Ln\textsuperscript{3+} and SiO\textsuperscript{−} are 3.5-4.5 Å apart, from Ln\textsuperscript{3+} completely dissociated from SiO\textsuperscript{−}.\textsuperscript{49} Furthermore, for Lu\textsuperscript{3+}, \( N_{hyd} \) does not increase monotonically as the cation desorbs.

Another, weaker harmonic potential of the form \( B_o[(\delta x - x_o)^2 + (\delta y - y_o)^2] \) constrains the Ln-O distances in the lateral directions. Here \( B_o=0.025 \) eV, \( \delta x=x_{Ln} - x_O, x_o \) is the equilibrium value of \( \delta x \) computed in completely unconstrained AIMD trajectories, and \( y_o \) and \( y_o \) are defined in analogous ways.

\( \Delta W(Z) \) is effectively the constrained free energy at a \( Z \) value; it does not include the standard state reference associated with aqueous solutions. To obtain the adsorption free energy \( (\Delta G_{ads}) \) from \( \Delta W(Z) \), we integrate configuration space in three dimensions, and account for the entropic contribution from a standard state 1.0 M ideal concentration solution:\textsuperscript{50}

\[
\Delta G_{ads}/k_BT = -\log \left\{ \int_{\Omega} d\Omega \exp[-\Delta W(Z)/k_BT]/(V_o) \right\}.
\] (1)

Here \( V_o \) is the volume associated with 1.0 M aqueous solution (1662 Å\textsuperscript{3}) and \( T=300 \) K is assumed. (“Standard state” refers to [Ln\textsuperscript{3+}]=1.0 M; no attempt is made to adjust the pH in Eq. (1).) The volume element \( \Omega \) spans the configuration space where Ln\textsuperscript{3+} is “bonded” to the SiO\textsuperscript{−} group. A limiting bonding distance of 3.20 Å is assumed. At this separation the pair correlation functions between transition metal ions and water oxygen sites exhibit their first correlation minima (Fig. S1 in the ESI). The angular distribution is also involved in the integral. To our knowledge, \( \Omega \) has not been standardized for PMF calculations at interfaces.\textsuperscript{48,49,51} Here we approximate it as a cylinder with a radius \( R=0.5 \) Å. Electrostatic corrections associated with image dipoles are added to the PMF predictions by creating a lattice model with screened coulomb interactions.\textsuperscript{25}

We do not apply the metadyamics method, based on non-equilibrium trajectories, to compute the PMF.\textsuperscript{52} The umbrella sampling approach used herein permits us to run trajectories of variable lengths not determined ahead of time. We examine statistical uncertainties
in each window to make sure there is no large, systematic drift in $\Delta W(R)$ in each AIMD trajectory.

III. RESULTS AND DISCUSSIONS

A. Batch Adsorption

First we discuss batch adsorption results on amorphous silica surfaces that motivated this work (Fig. 1). The experiments were performed on silica surfaces with $\sim$2 SiOH groups per nm$^2$ of surface, as discussed in our earlier publication. The lanthanide adsorption data was fit using the Langmuir isotherm model. Based on the fitting, the maximum adsorption coverage for Eu$^{3+}$ was estimated at 20.4 $\mu$moles/g, and the Langmuir constant $K_L$ was 0.102 L/g. For Lu$^{3+}$ the maximum adsorption was estimated at 19.6 $\mu$moles/g, and $K_L$ at 0.138 L/g. The overall affinity at pH 5.0 for Lu$^{3+}$ and Eu$^{3+}$ was similar, with Lu$^{3+}$ being slightly more favorable. This suggests that the experimental $\Delta G_{ads}$ for Eu$^{3+}$ and Lu$^{3+}$ are very similar.

B. Potential-of-Mean-Force

Next we turn to AIMD modeling. Fig. 2a depicts the charge-neutral simulation cell containing 123 H$_2$O molecules and a partially deprotonated, surface-reconstructed $\beta$-cristobalite slab; Lu$^{3+}$ is coordinated to two SiO$^-$ groups on one surface. Fig. 2b-e depict Lu$^{3+}$ con-
FIG. 2: (a) The 14.3×14.3×26 Å³ simulation cell with adsorbed Lu³⁺ complex coordinated to two SiO⁻ groups. The reaction coordinate Z is along the z direction (left-to-right). (b)-(c) Z centered at \( Z_i = 4.9 \) Å, with Lu³⁺ bound to two or one SiO⁻ groups. (d) \( Z_i = 7.4 \) Å. (e) \( Z_i = 8.0 \) Å. \( Z_i \) is the center of the constrained window. Si, O, O (water), H, and Lu are depicted in yellow, red, blue, white, and pink. As some protons in H₂O are obscured, the true OH⁻ species are circled in red.

Fig. 3a compares \( \Delta W(Z) \) for Lu³⁺ and Eu³⁺. The shapes of \( \Delta W(Z) \) at small \( Z \) are similar, suggesting similar energetics in the neighborhood of the optimal adsorption configuration. As desorption proceeds and \( Z \) approaches \( Z \sim 5 \) Å, a cross-over to a quasi-plateau region is observed. The Eu³⁺ potential-of-mean-force exhibits a slight repulsive behavior (\( \Delta W(Z) > 0 \)) near \( Z = 5 \) Å. This is likely related to overscreening behavior associated with multivalent electrolytes.\[^{53}\] In contrast, due to hydrolysis in its hydration shell (see below), the more weakly charged, hydrolyzed Lu³⁺(OH⁻)_n complexes have a lower net charge and a monotonic \( \Delta W(Z) \). Our AIMD simulations are not ideally suited to investigating overscreening effects due to the lack of counter-ions. However, our future classical force field MD simulations will reconsider possible overscreening. Neither cation exhibits a local minimum.
associated with outer sphere solvation; local minima may have been helpful in engineering preferential adsorption motifs.

Integrating $\Delta W(Z)$ yields $\Delta G_{\text{ads}} = -0.79 \pm 0.04$ eV and $-0.84 \pm 0.03$ eV, assuming standard states for Ln$^{3+}$ and Eu$^{3+}$. The uncertainties reflect one standard deviation. Not included in Fig. 3 are electrostatic corrections. As the Ln$^{3+}$ desorbs, a significant dipole moment is created in the simulation cell, leading to image-image repulsions in the lateral directions. Using corrections based on lattice-models with dielectric screening the magnitudes of $\Delta G_{\text{ads}}$ are reduced by 0.01 and 0.09 eV for Ln$^{3+}$ and Eu$^{3+}$, respectively. These corrections change the preferred adsorption from Eu$^{3+}$ over Ln$^{3+}$ to Ln$^{3+}$ over Eu$^{3+}$ by 0.03 eV, because the Lu$^{3+}(\text{OH}^-)_n$ complex incurs less correction. The computed $\Delta G_{\text{ads}}$ are larger in magnitude than our AIMD predictions of divalent metal cations on mineral surfaces, which range from -0.38 to -0.71 eV. Our predicted Ln$^{3+}$ $\Delta G_{\text{ads}}$ are more negative than those measured for some trivalent cations on silica surfaces, likely because of the higher effective pH in our simulations, our absence of counter-ions, and possible differences in SiOH spatial distributions between our model and the experiment samples. The measurements in Ref. 18 are conducted at pH=4 while the pH in our simulation cell is estimated to be $\sim 7.5$. A 3.5 unit increase in pH translates into a maximum of $2 \times 3.5 \times 0.059 \text{ eV} = 0.41$ eV increase in Ln$^{3+}$ binding free energy. This estimate assumes that the increase in pH reduces the free energy needed to deprotonate two neighboring SiOH groups, which then bind to the Ln$^{3+}$. It is much more difficult to quantify the dependence of $\Delta G_{\text{ads}}$ on surface structure details, such as the distance between the two SiO$^-$ groups coordinated to the adsorbed Ln$^{3+}$, without explicit AIMD/PMF simulations of the modified structure. We plan to pursue AIMD simulations of monodentate Ln$^{3+}$ on model silica surfaces in the future.

As discussed in Sec. II C, around $Z_i = 5.0 \text{ Å}$, a secondary PMF calculation, with another reaction coordinate $R'$, which is the true distance between the Ln$^{3+}$ and a flagged O atom (not just its $z$-component), is needed to augment our results. Fig. 4a shows that the Eu$^{3+}$ $\Delta W(Z)$ in the $Z<5.0 \text{ Å}$ and $Z>5.0 \text{ Å}$, windows (green and blue lines) have different slopes. Combining these curves would yield a sharp kink in $\Delta W(Z)$. A similar kink would have occurred in the Lu$^{3+}$ $\Delta W(Z)$ (Fig. 4c). These kinks signal the inability of AIMD/PMF simulations to reversibly sample two free energy valleys separated by small barriers. Fig. 2b-c show that these valleys are in fact associated with Lu$^{3+}$ coordinated to one and zero SiO$^-$ groups, respectively; the local minima are separated by Lu$^{3+}$ displacement parallel to the silica surface. In terms of the secondary coordinate, $R'$ jumps from 2.5 to 4.2 Å between these valleys. Our secondary PMF $\Delta W'(R')$ estimates the free energy differences between these two valleys (Sec. II C) and largely smooths over the kink in $\Delta W(Z)$. The barrier between the valleys are 0.2 to 0.25 eV (Fig. 4b,d), which are small but would have required much longer AIMD trajectories to sample adequately without the secondary PMF. While this approach involves approximations (Sec. II C), the uncertainty in $\Delta \Delta G_{\text{ads}}$ is lessened
C. Surprisingly Large Energy Difference in Adsorbed States

The predicted preferential Lu\(^{3+}\) adsorption is qualitatively consistent with our batch adsorption measurements (Fig. 1). The difference between the Lu\(^{3+}\) and Eu\(^{3+}\) \(\Delta G_{\text{ads}}\) is small, comparable to the statistical uncertainty. However, this small \(\Delta \Delta G_{\text{ads}}=-0.03\) eV, is surprising from an energetic standpoint – despite the much-quoted lanthanide “chemical similarity.” As mentioned above, Lu\(^{3+}\) exhibits \(\Delta G_{\text{hyd}}\) which is more favorable (negative) than the Eu\(^{3+}\) value by -2.17 eV.\(^{16}\) This represents the desorption end point behavior. For the two \(\Delta G_{\text{ads}}\) to be similar, there must be a similarly large energetic difference at the adsorbed end point where Lu\(^{3+}\) and Eu\(^{3+}\) are in contact with silica. In other words, chemical similarity in fact derives from a cancellation of large (relative) energy terms.

To examine this hypothesis, we approximate the energy difference (\(\Delta \Delta E_{\text{ads}}(\text{dry})\)) in the adsorbed state by omitting most water molecules. We optimize configurations with a Lu\(^{3+}\) or Eu\(^{3+}\) cation at the binding site coordinated to two SiO\(^-\) groups at T=0 K (Fig. 5a-b). Only one H\(_2\)O molecule is included in the simulation cell. Maximally localized Wannier
FIG. 4: Panels (a)-(b) and (c)-(d) refer to $\Delta W(Z)$ for Eu\textsuperscript{3+} and Lu\textsuperscript{3+}, with $Z_i=5.0\ \text{Å}$ or $Z_i=4.9\ \text{Å}$, respectively. Green and blue are unnormalized $\Delta W(Z)$ segments in the two $R'$ free energy valleys with the same $Z_i$ constraint; they correspond to Ln\textsuperscript{3+} coordinated to one or zero SiO\textsuperscript{−}. Red depicts a weighted average of the two, based on secondary umbrella sampling. (b) and (d): potential-of-mean force along a secondary reaction coordinate $R'$ ($\Delta W(R')$), with $R'$ being one of the O-Ln distances. See text.

Function analysis\textsuperscript{54} confirms that, in these charge-neutral simulation cells with significant vacuum regions, both lanthanides remain trivalent cations. The net energy of the Lu\textsuperscript{3+} simulation cell is lower than that of Eu\textsuperscript{3+} by $\Delta\Delta E_{ads}(\text{dry})=-1.68\ \text{eV}$ after substracting the respective gas phase, bare ion energies. This difference is indeed similar to the reported -2.17 eV difference in $\Delta E_{\text{hyd}}$.\textsuperscript{15} Note that we cannot report accurate absolute binding energies between Ln\textsuperscript{3+} and negatively charged silica because of the difficulty in correcting the energies of slabs with net charges.
FIG. 5: (a)-(b) Top and side views of optimized Lu$^{3+}$ adsorption configuration on silica surface, at $T=0$ K and coordinated to one H$_2$O molecule. (c)-(d) Similar to (b) but with two and three H$_2$O coordinated to Eu$^{3+}$, respectively.

D. Justification for Using the Eu(A) Pseudopotential

The systems depicted in Fig. 5 also represent convenient platforms to examine the validity of the Eu(A) pseudopotential used, which omits $f$-electrons. It is tempting to assume that Eu(B), which includes $f$-electrons and requires the spin-polarized DFT method (Sec. II B), would be more accurate. Our attempts at using Eu(B) in DFT/PBE-based AIMD simulations, however, result in occasional failures in the self-consistent field procedure when performing convergence of the Kohn-Sham densities and the Slater determinant orbitals. The likely reason is an unphysical charge transfer from silica to the Eu$^{3+}$ when using this pseudopotential during AIMD. This is likely a failure of the PBE functional, which is known to predict unphysical electron delocalization in $f$-electron systems like CeO$_2$.\footnote{55,56} To deal with this problem, the DFT+U method\footnote{36} has often been applied as a remedy,\footnote{57} as have hybrid DFT functionals like HSE06.\footnote{37,38}

Here we compute the Eu(B)-predicted binding energies of a Eu$^{3+}$ relative to Lu$^{3+}$ to the Si$_{40}$O$_{88}$H$_{13}^{-}$ slab (Fig. 5b) in vacuum, using the Eu(A) value as a reference. If both Eu pseudopotentials are equally accurate within the DFT/PBE framework, the energy difference ($\Delta\Delta E$) between them should be zero. Instead, we find that the Eu(B) result is favored by
\( \Delta \Delta E = -0.70 \text{ eV} \) over Eu(A). When two or three \( \text{H}_2\text{O} \) are included (Fig. 5c-d), \( \Delta \Delta E = -0.51 \text{ eV} \) and -0.48 eV, respectively.

Eu(B) gives consistently lower energies. We argue that the significantly more negative \( \Delta \Delta E \) is consistent with unphysical hybridization between silica and Eu(B) 4\( f \)-orbitals, and/or possible electron transfer from silica to Eu\( ^{3+} \). Note that the extent of charge-transfer is non-trivial to quantify; Wannier function analysis is challenging when using Eu(B), because the partially-filled \( f \) shell gives “metallic” behavior. To support our argument, we turn to a rotationally invariant DFT+U approach\(^ {36} \) with \( U-J = 4.5 \text{ eV} \). We find that \( \Delta \Delta E = -0.13 \text{ eV}, -0.11 \text{ eV}, \) and -0.04 eV with 1-3 \( \text{H}_2\text{O} \) in the simulation cell. Therefore the Eu(A) pseudopotential yields predictions very similar to Eu(B) which has \( f \)-electrons – as long as the more reliable\(^ {57} \) DFT+U augmentation is applied to \( f \)-electrons in the latter case. Eu(A) does not have 4\( f \) electrons and DFT+U is inapplicable there, while Lu has a full 4\( f \) shell and DFT+U is not expected to yield results significantly different from PBE predictions. We also note that the Eu(A) and Eu(B) pseudopotentials have been shown to yield similar structural properties when the latter is used in conjunction with DFT+U augmentation.\(^ {58} \)

Although applying DFT+U alongside the Eu(B) pseudopotential gives better agreement with Eu(A), the results slightly vary with the value of \( (U-J) \). Sec. S3 in the SEI reports that changing \( (U-J) \) from 4.5 eV to 6.5 eV for the one-water (Fig. 5h-b) configuration changes \( \Delta \Delta E \) from -0.13 eV to 0.05 eV. Although this variation is only 0.18 eV, it is significant compared with the \( \Delta \Delta G_{\text{ads}} = 0.03 \text{ eV} \) difference computed in AIMD simulations. To lessen this ambiguity, we also apply the HSE06 functional to both Lu\( ^{3+} \) and Eu\( ^{3+} \) bound to silica surfaces. The HSE06 \( \Delta \Delta E \) for the Fig. 5h-b configuration is found to be -0.06 eV, which is close to zero. We argue that the Eu(A) pseudopotential, used in the majority of this paper, is an approximate way to implement HSE06, or DFT+U with \( U-J = 4.5 \text{ eV} \), on the Eu pseudopotential with \( f \)-electrons. It should in fact give more physical results than Eu(B) when the latter is applied with PBE only. From these state calculations, we estimate that the systematic uncertainty associated with using the Eu(A) pseudopotential in our AIMD \( \Delta \Delta G_{\text{ads}} \) calculations is between \( \sim 0.06 \) and 0.13 eV.

E. Hydrolysis and Hydration Behavior

In the rest of this paper we analyze the differences in Lu\( ^{3+} \) and Eu\( ^{3+} \) hydrolysis and hydration properties in an attempt to identify factors that promote preferential Ln\( ^{3+} \) adsorption and separation. As noted above, initially Lu\( ^{3+} \) is coordinated to two SiO\(^- \) groups and 4 H\( \text{H}_2\text{O} \) molecules (Fig. 2h). Bidentate adsorption is consistent with dihydroxyl Yb\( ^{3+} \) coordination at low Yb\( ^{3+} \) coverage known from previous analysis.\(^ {59} \) In the cross-over region \( (Z \sim 4.9 \text{ Å}) \), there is an equilibrium between two states, with Lu\( ^{3+} \) coordinated to one (Fig. 2b) and zero SiO\(^- \) (Fig. 2c) group. Hydrolysis only occurs above \( Z \sim 5 \text{ Å} \); it arises
from the transfer of proton(s) from H₂O molecule(s) coordinated to the Lu³⁺, to the silica surface via the Grothuss mechanism. When the desorption is almost complete (Fig. 21-e), Lu³⁺ forms a Lu³⁺(H₂O)₆(OH⁻), a Lu³⁺(H₂O)₅(OH⁻)₂, or a Lu³⁺(H₂O)₄(OH⁻)₂ complex.

Eu³⁺ configurations (not shown) are qualitatively similar. Two main differences are that Eu³⁺ exhibits less tendency towards hydrolysis than Lu³⁺, and has larger hydration numbers (N_{hyd}) throughout the entire Z range. First we focus on hydrolysis. Fig. 3b reports N_{sio} as Z varies. N_{sio} counts the number of deprotonated Si-OH surface groups with a 1.25 Å O-H cutoff distance in each sampling window each centered at Z=Z_i. N_{sio} plus N_{oh} (the number of hydrolysis events or number of OH⁻ coordinated to Ln³⁺) should add to 3.0 on average. When Z>5 Å, the average N_{sio} is larger in Eu³⁺ simulation cells compared to Lu³⁺, meaning Eu³⁺ induces less hydrolysis. Like Lu³⁺, the onset in SiO⁻ protonation state change is correlated with the onset of the Eu³⁺ ∆W(Z) plateau.

Based on the above analysis, it is surprising that a substantial cancellation in energetics occurs for these two cations during desorption. Although the Lu³⁺/SiO₂ (Fig. 5i) and the Eu³⁺/SiO₂ binding configurations are similar (the latter is not shown), Lu³⁺ is coordinated to OH⁻ while Eu³⁺ is coordinated only to H₂O far from the surface. We conjecture that a substantial energy cancellation may arise from the fact that the pK_{a₁} for hydrolysis is not far from the simulation cell pH condition, so there is little change in ∆∆G_{hyd} regardless of whether hydrolysis occurs. Indeed, beyond Z=5 Å, ∆W(Z) only changes by ~0.1 eV in the ∆W(R) for both cations. Hence we argue that hydrolysis does not strongly affect the predicted ∆∆G_{ads} for either Eu³⁺ or Lu³⁺. This is unlike the case of Cu²⁺. We propose that, if a local pH significantly higher than the Ln³⁺(H₂O)ₙ pK_{a₁} can be maintained, this crucial cancellation can be disrupted without causing Ln(OH)₃ precipitation, leading to more selective Ln³⁺ adsorption on silica surfaces.

Next we examine how differences in hydration numbers can affect ∆∆G_{ads}. Fig. 3b also reports N_{hyd}, which is the number of H₂O or OH⁻ oxygen atoms within 3.2 Å of Ln³⁺. It does not count coordination to SiO⁻ groups. Eu³⁺ exhibits hydration numbers which exceed those of Lu³⁺ by about 1-2 in the entire Z range, i.e., ∆N_{hyd}~1-2. Note that Ln³⁺ hydration numbers in liquid water are measurable in X-ray and neutron scattering experiments, and much computational effort has been devoted to reproducing those values. We do not observe Lu³⁺(H₂O)₈ and Eu³⁺(H₂O)₉ complexes which have been predicted in quantum chemistry or molecular dynamics simulations. One reason is the Lu³⁺ inner sphere hydrolysis behavior discussed above, which has not been accounted for in previous modeling work. Another reason may be that the explicit treatment of the second hydration shell in AIMD simulations changes the first shell hydration number computed using an implicit solvent approximation in the literature. Finally, different DFT functionals can yield N_{hyd} which are slightly different from each other and from experiments. The free energy difference associated with N_{hyd} values that differ by one is generally on the order
of $k_B T = 0.025$ eV, which is small on our $\Delta W(Z)$ energy scale. We propose that surface constraints or functional groups that increase $\Delta N_{\text{hyd}}$ to 2 or higher may be needed to aid selective adsorption.

F. Sampling Dynamics

For completeness, we briefly discuss dynamics. $N_{\text{hyd}}$ and $N_{\text{oh}}$ for Lu$^{3+}$ are depicted as functions of time in selected sampling windows in Fig. 6. (The complete set is given in the ESI.) We have plotted $N_{\text{oh}}$ instead of $N_{\text{sio}}$ here because, at certain times, a H$^+$ may be in transit from the hydration shell of the desorbed Ln$^{3+}$ to the silica surface; therefore $N_{\text{oh}}(t)$ is more descriptive than $N_{\text{sio}}$ which equals $3-N_{\text{oh}}$ only on average. At $Z>5$ Å, the average $N_{\text{oh}}$ is not monotonic as $Z$ varies, and there are picosecond time scale proton exchanges between the Lu$^{3+}$ inner hydration shell and the silica surface not accompanied by significant changes in $\Delta W(Z)$ (Fig. 6). We never observe Si-OH groups from the opposite surface of the silica slab being involved in acid-base reactions.

Fluctuations of $N_{\text{hyd}}$ for trivalent cations can take long times at room temperature. Fortunately, our sampling efficiency is improved by a slightly elevated temperature of 400 K. The fast proton transfer dynamics and the large driving force guiding $N_{\text{hyd}}$ evolution as $Z$ varies also help inner sphere equilibration. As a test, we have confirmed that, after removing the umbrella sampling constraint for Lu$^{3+}$ at $Z_i=3.5$ Å, $N_{\text{hyd}}$ spontaneously relaxes from 5 to its equilibrium value of 4 (Fig. 3b) within 5 ps as the Lu$^{3+}$ relaxes to its adsorbed configuration. See also the ESI for discussions of the special case of the $N_{\text{hyd}}$ in one Lu$^{3+}$ window. However, our simulation time scales would have been insufficient to deal with the presence of anions, which diffuse through water much more slowly than H$^+$. This is the main reason we have not included explicit counter anions in our AIMD simulations, despite the fact that our batch adsorption experiments include NO$_3^-$.

IV. CONCLUSION

In summary, using AIMD/PMF calculations, we predict a slight, 0.03±0.05 eV preferential adsorption of Lu$^{3+}$ over the lighter, larger Eu$^{3+}$ on model silica surface with deprotonated silanol groups at its interface. This finding is in qualitative agreement with our batch adsorption work. The 0.03 eV difference should be considered qualitative because of differences between the experimental and computational systems, which include silica surface details; pH; the presence/absence of NO$_3^-$ counter-ions; multiple surface binding sites; and possible Lu$^{3+}$ dimerization. We also note that our Lu$^{3+}$ and Eu$^{3+}$ calculations utilize Lu and Eu pseudopotentials with and without explicit 4$f$-electrons, respectively. As a result, the Lu$^{3+}$ predictions are likely more accurate than the Eu$^{3+}$ predictions. It is challenging to model
Ln$^{3+}$ in aqueous media, both because of DFT functional accuracy issues and because of the statistical mechanical details that need to be implemented to obtain accurate free energy changes associated with highly charged trivalent cations which can desorb in conjunction with hydrolysis reactions. In that sense, our pioneering AIMD simulations pave the way for future examination of these details via systematic variation of the model surface. We explicitly address the role of concerted proton motion, and give quantitative desorption free energy predictions for trivalent lanthanide cation adsorption on mineral surfaces. Our predictions provide guidance to molecular dynamics simulations that apply classical force fields.

AIMD simulations are first-principles in nature, and are generally more accurate than force field-based MD. They are computationally costly; the necessarily limited trajectory
lengths lead to unavoidable statistical uncertainties. A \( \sim 10 \)-fold preference for Lu\(^{3+} \) over Eu\(^{3+} \) at \( T=300 \) K, measured under conditions slightly different from this work,\(^{22} \) translates into a -0.059 eV difference in \( \Delta G_{\text{ads}} \), which is almost within AIMD noise level. Instead of resolving small \( \Delta G_{\text{ads}} \) differences, AIMD is most valuable at providing mechanistic insights. Thus it is more significant that the similarity of the Lu\(^{3+} \) and Eu\(^{3+} \) desorption free energies is found to arise from a cancellation between their relative adsorption energies to deproto- nated silica surface (\( \Delta \Delta E_{\text{ads}}(\text{dry}) \)), and their relative hydration free energies in liquid water (\( \Delta \Delta G_{\text{hyd}} \)). Although the cations exhibit substantial differences in hydration and hydrolysis properties, the cancellation leads to very similar \( \Delta G_{\text{ads}} \). To disrupt this cancellation of energy differences, enhance Ln\(^{3+} \) selectivity, and aid separation, we propose that modification of the silica surface to change the local pH or hydration environment, would be valuable. Another approach suggested by our analysis is to modify \( \Delta G_{\text{hyd}} \) in the desorbed state, \( e.g., \) by using a mixed solvent, or \textit{via} nanoconfinement.\(^{22} \)

**Conflicts of Interest**

Nothing to declare.

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