Exponential Scaling of Water Exchange Rates with Ion Interaction Strength from the Perspective of Dynamic Facilitation Theory

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Water exchange reactions around ionic solutes are ubiquitous in aqueous solution-phase chemistry. However, the extreme sensitivity of exchange rates to perturbations in the chemistry of an ionic solute is not well understood. We examine water exchange around model ions within the language of dynamic facilitation theory, typically used to describe glassy and other systems with collective, facilitated dynamics. Through the development of a coarse-grained, kinetically-constrained lattice model of water exchange, we show that the timescale for water exchange scales exponentially with the strength of the solute-solvent interactions.

I. INTRODUCTION

Changes in coordination structures, particularly of coordinated water molecules, are fundamental reactions of general importance in chemistry. Understanding the molecular mechanisms underlying their chemical kinetics is an important step in fully characterizing more complex reactions of broad importance. For example, mineral dissolution and precipitation in aqueous solution involves the coordination (and de-coordination) of metal ions and their complexes by water molecules, and water exchange has been suggested to be a rate-limiting step in dilute solutions. In biological settings, water exchange plays an important role in the binding of ions to RNA and protein, influencing rates of biological processes. Additionally, coordination complexes involving mixtures of water and a cosolvent or pure non-aqueous solvents have proved important to the development of complex materials for catalysis and energy storage.

The exchange of a hydration water for another around an ion (M) — the water exchange reaction — is one of the (seemingly) simplest changes in coordination structure, where $^{*}$OH$_2$ is used to identify a distinct water molecule, and the ion M has a coordination of n in water. The reactants and products are chemically identical in this process, such that there is no net change in the free energy driving this reaction. However, the kinetics governing water exchange are quite complex. Typical timescales for water exchange depend on the nature of the ion and vary from picoseconds to hundreds of years.

In this work, we examine the complex dynamics of water exchange reactions within the framework of dynamic facilitation (DF) theory, typically employed to describe glassy systems. To do so, we suggest that the coordination of a solute by a water molecule imposes a kinetic constraint on the dynamics of that molecule. We additionally suggest that these kinetic constraints are coupled to the charge density of the solute, and, in general, any solute-solvent interactions, such that strong solute-solvent interactions ultimately lead to slow, collective dynamics that are dominated by fluctuations, precisely the realm of applicability of DF theory. We use computer simulations to characterize water exchange around a range of model ions within the DF theory framework, illustrating that increased water exchange times correspond to increased glassiness and collective behavior in the ion hydration shell. Furthermore, by using DF theory and developing kinetically constrained lattice models, we predict that water exchange times scale exponentially with the strength of water-solute interactions, in agreement with recent empirical findings.

II. SIMULATION DETAILS

We probe the dynamics of water exchange around classical (Lennard-Jones plus point charge) ion models using GROMACS version 5.0. We do not consider the influence of changes in electronic structure, bond breaking, and nuclear quantum effects, which may impact water exchange kinetics. All simulations were performed in the isothermal-isobaric (NPT) ensemble at a temperature of $T = 300$ K and a pressure of 1 bar using the canonical velocity rescaling thermostat and an Andersen barostat, respectively. Water was modeled using the extended simple point charge (SPC/E) model. We study several model ions to span a significant range of water exchange times: (i) Ca$^{2+}$, (ii) a model Cobalt ion with a reduced charge of 1.4 (Co$^{1.4+}$) in order to make direct sampling of water exchange possible, and (iii) an model potassium ion (K$^+$. Co$^{1.4+}$ was modeled using the CM parameters developed by Merz and coworkers, and the LJ parameters of the ECCR model were used for Ca$^{2+}$, with a charge of +2, neglecting any charge transfer between the ion and water. K$^+$ was modeled using the parameters developed by Konshan et al. Short-ranged electrostatic and Lennard-Jones (LJ) interactions were truncated at a distance of 1 nm. The particle mesh Ewald method was used to evaluate
long-ranged electrostatic interactions with the inclusion of a uniform neutralizing background potential to cancel the net charge of the single ion in the system. Equations of motion were integrated using a timestep of 0.5 fs, saving configurations every 100 timesteps, employing the LINCS algorithm to fix the O-H bonds and H-O-H angles of water. Trajectory lengths for $\text{Co}^{1.4+}$, $\text{Ca}^{2+}$, and $\text{K}^+$ systems were 1 μs, 711 ns, and 100 ns, respectively. Unless otherwise noted, results are presented for systems with 500 ($\text{Co}^{1.4+}$ and $\text{K}^+$) or 512 ($\text{Ca}^{2+}$) water molecules, which are large enough to minimize finite size effects (see Supporting Information (SI)).

III. FACILITATED DYNAMICS OF WATER EXCHANGE

In order to quantify the kinetics of water exchange, we must first define what we mean by the exchange of a water molecule. Following previous work, we monitor exchange through the indicator function $h_i(t)$, which is equal to one if water $i$ is in the hydration shell of an ion and is equal to zero when it is not. A water molecule is in the first coordination shell if the ion-water oxygen distance is less than $r_c$, defined as the position of the free energy barrier in the solute-water oxygen potential of mean force, $\beta W(r) = -\ln g(r)$, shown in Fig. 1a for the ions under study, where $g(r)$ is the water oxygen-ion pair distribution function, which are shown and discussed in the SI, along with the ion coordination structure. The free energy barriers increase following $K^+ < \text{Ca}^{2+} < \text{Co}^{1.4+}$, and we therefore expect the water exchange times to follow a similar trend.

Water exchange times are traditionally measured by computing the time correlation function

$$C(t) = \frac{\langle \delta h_i(0) \delta h_i(t) \rangle}{\langle \delta h_i^2(0) \rangle}, \quad (2)$$

which is shown in Fig. 1b, where $\delta h_i(t) = h_i(t) - \langle h_i(t) \rangle$ and implicit in the ensemble average is an average over all water molecules. For tightly bound water molecules, as is the case for $\text{Ca}^{2+}$ and $\text{Co}^{1.4+}$ model ions, the decay of $C(t)$ is exponential, yielding water exchange lifetimes of roughly 200 ps for $\text{Ca}^{2+}$ and 2000 ps for the reduced charge $\text{Co}^{1.4+}$ model, both of which are in good agreement with earlier estimates for similar model ion parameters. Note that water exchange dynamics can display significant finite size effects, as discussed in the SIs for $\text{Ca}^{2+}$. For $K^+$, $C(t)$ decays exponentially only after an initial transient period. This transient short-time dynamics physically arises from high-frequency recrossing of the barrier at $r_c$ and is removed upon coarse-graining in the time domain, as is typically done for glassy systems. The coarse-grained $C(t)$ for $K^+$ decays exponentially with a time constant of roughly 10 ps, where the coarse-graining is performed over a time window of 2 ps. Coarse-graining the correlation functions for $\text{Ca}^{2+}$ and $\text{Co}^{1.4+}$ on this time scale does not alter their behavior.

Exponential decay of correlation functions is typically indicative of uncorrelated dynamics, not collective, heterogeneous dynamics. However, we show below that the dynamics of water exchange are facilitated and collective, despite the exponentially decaying $C(t)$. Instead, we suggest that this exponential decay arises from the small phase space volume probed by these correlation functions.

We characterize the facilitated nature of the dynamics of water exchange by computing the probability distributions of exchange and persistence times, $P_X(t)$ and $P_t(t)$, respectively, using the language of DF theory. The
exchange time is defined as the time between two successive events. In contrast, the persistence time is the time for the first event to occur, as measured from $t = 0$. The distributions of exchange and persistence times are related through

$$P_P(t) \sim \int_0^\infty dt' P_X(t'),$$

within a normalization constant.

We define an event as any change in the water coordination shell between two successive timesteps, mathematically determined by the indicator function

$$I(t) = \Theta(\Delta H(t) - c),$$

where $\Theta(x)$ is the Heaviside step function, $c$ is a constant greater than zero, $\Delta H(t) = \sum_{i=1}^N \Delta h_i(t)$, and $\Delta h_i(t) = |h_i(t) - h_i(t - \Delta t)|$. The constant $c$ sets the threshold for defining an event when the trajectory is coarse-grained in time to avoid counting transient barrier recrossing, and here we choose $c = 0.9$ without impact on the qualitative findings. A representative portion of an $I(t)$ trajectory is shown in Fig. 2, with individual exchange and persistence times highlighted. This trajectory suggests that dynamics of water exchange are indeed facilitated. One observes long periods of inactivity and bursts of activity that are localized in time. This complex dynamic structure is quantified by the exchange and persistence time distributions.

For a Poisson process, the distribution of bound water lifetimes is exponential, and the persistence and exchange time distributions coincide and are exponential.

In the context of facilitated dynamics, a dynamical event cannot occur unless a molecule is intersected by dynamical excitations in spacetime, which become increasingly sparse under conditions where the dynamics are slowed. In supercooled liquids, the concentration of excitations is typically controlled by the temperature. In the context of water exchange, the relevant excitations are those that permeate the spacetime of the hydration shell, and the density of these excitations is expected to be inversely related to the solute-solvent interaction strength, in addition to the temperature of the solution. In the limit of high solute charge density, for example, the excitations will be sparse, leading to persistence times that are significantly decoupled from exchange times. This behavior is exemplified by the exchange and persistence time distributions in Fig. 2.

As the solute charge density is increased, spacetime excitations in the hydration shell become increasingly...
sparse, and there is a small number of dynamic pathways that enable hydration water to exchange. Consequently, the exchange and persistence time distributions become increasingly decoupled as the solute-solvent interaction strength is increased, Fig. 2a-c. This illustrates that strong solute-solvent interactions lead to facilitated and collective water exchange processes. For weak solute-solvent interactions (Fig. 2d), as is the case for K⁺, water molecules in the hydration shell are not trapped by strong solute-solvent interactions, and the barrier to water exchange is small (Fig. 1h). In this case, excitations permeate the spacetime of the solute hydration shell, there is a high density of dynamic pathways that enable water exchange to occur, and the exchange and persistence time distributions coincide. We expect that solutes with net repulsions, including effective repulsions leading to drying around large hydrophobic solutes, will also lead to \( P_X(t) \) and \( P_Y(t) \) that coincide.

We also note that the exponential decay of \( C(t) \) shown in Fig. 1h, which seems to contradict the signatures of dynamic heterogeneity found here, is expected to arise from the small spacetime volume in which the system is kinetically constrained. We expect that a stretched exponential (0 → 1) occurring at a rate of \( c \) will also lead to \( P_X(t) \) and \( P_Y(t) \) that coincide.

We illustrate this within the context of the Fredrickson-Andersen (FA) model, but our approach is general and can be readily extended to other kinetically constrained models, such as the East model. In the FA model, each lattice site is described by a binary variable \( n_i \) = 0, 1, typically interpreted as (0) immobile and (1) mobile, or excited, sites. The energy of a FA model with \( N \) sites is \( E = J \sum_{i=1}^{N} n_i \), where \( J \) is an energy scale. The resulting thermodynamic properties of the FA model are trivial. In particular, the concentration of excitations (mobile sites) is \( c_{FA} = \langle n_i \rangle = 1/(1 + e^{1/\tilde{T}}) \), where \( \tilde{T} = k_B T / J \). The complex dynamics of the FA model arise from kinetic constraints imposed on the system. The kinetic constraints dictate that a change in state of a site can only occur if at least one of its neighboring sites is excited, with excitations (0 → 1) occurring at a rate of \( c_{FA} \) and de-excitations (1 → 0) occurring at a rate of \( 1 - c_{FA} \).

In the water exchange reaction, the solute imposes kinetic constraints largely on the water molecules in its coordination shell only, not the rest of the solvent. Thus, we introduce a fixed solute site into the lattice model, which, for simplicity, does not move or change state. In order to mimic the water exchange reaction, we impose the FA kinetic constraints only on the nearest neighbor sites of the solute; the remaining lattice sites exhibit unconstrained dynamics. The solute is considered to be an immobile region; it does not facilitate the dynamics of its neighbors.

Finally, we note that the kinetics of the ionic solvation shell water molecules are altered by the nature of the solute, through solute-solvent interactions. To mimic this, we assign a different energy scale, \( J' \), to the nearest neighbors of the solute in the lattice model, arising physically from any solute-solvent interactions present. Defining \( \tilde{T}' = k_B T / J' \), one can consider the dynamics of these sites to evolve at an effective reduced temperature \( \tilde{T}' \), which is different than that of the rest of the system, \( \tilde{T} \). In this context, \( \tilde{T}' \) captures the effects of the solute charge density and any other solute-solvent interactions (e.g. LJ interactions). For example, lowering \( \tilde{T}' \) is the analog of increasing the charge density of the ion.

Here, we focus on the one-dimensional (1D) version of this model. The thermodynamics of this model are trivial and analogous to that of the FA and East models. For \( S \) solutes, dispersed far apart (their solvation shells are not shared), there will be \( 2S \) solute neighbors in a 1D system. Omitting the (static) solutes from consideration, the (solvent) partition function is readily evaluated as

\[
\mathcal{Z} = (1 + e^{-1/\tilde{T}})^{N-2S} (1 + e^{-1/\tilde{T}'}^S)^{2S}.
\]

The concentration of excitations is obtained from the sum of the excitation concentrations in the constrained and...
unconstrained regions,

\[ c = \left(1 - \frac{2S}{N}\right) \frac{1}{1 + e^{1/T}} + \frac{2S}{N} \frac{1}{1 + e^{1/T'}} \]

or

\[ \equiv \left(1 - \frac{2S}{N}\right) c_B + \frac{2S}{N} c_{\text{CNN}}. \]  

(6)

The first term in Eq. (6) \( c_B \), corresponds to the concentration of excitations in the unconstrained, bulk (B) region of the system, i.e. the lattice sites that are not nearest neighbors of a solute. Correspondingly, the second term in Eq. (6) is the concentration of excitations in the regions composed of lattice sites that are nearest neighbors (NN) of a solute. Thus, Eq. (6) is equal to \( c_{FA} \) when \( S = 0 \) or \( T' = T \). Equation (6) illustrates that the role of the effective temperature \( T' \) is to alter the concentration of excitations in the solute solvation shell. Indeed, our interpretation of water exchange described in the previous section yields a similar picture for the effects of solute-solvent interactions, which are mimicked by \( T' \).

The corresponding distributions of the exchange and persistence times are decoupled for small \( T' \) (strong solute-solvent interactions), Fig. 3a-d, which were obtained from 100 independent trajectories at each \( T' \) using a 1D lattice with \( N = 50 \) and \( S = 1 \), evolving under Metropolis dynamics. As \( T' \) is increased, the exchange and persistence time distributions become increasingly similar. Physically, increasing \( T' \) corresponds to weakening the solute-solvent interactions, and even adding solute-solvent repulsions (\( T' > T \)), arising from direct or effective repulsions that can lead to drying around extended hydrophobic surfaces \( \text{[11][14][15]} \). Both reduce the kinetic constraints for water exchange and lead to overlapping distributions.

The average exchange and persistence times, \( \tau_X \) and \( \tau_P \), respectively, further support the decoupling of these two processes as \( T' \) is lowered, Fig. 4. For large values of \( T' \), \( \tau_X \) and \( \tau_P \) coincide. As \( T' \) is lowered, \( c_{\text{NN}} \) decreases, and \( \tau_X \) and \( \tau_P \) are increasingly decoupled. The dynamics controlling exchange and persistence in the solute nearest-neighbor lattice sites are facilitated by the unconstrained dynamics of the surrounding (bulk) lattice region, such that

\[ \tau_X \sim c_{\text{NN}}^{-1} \]  

and

\[ \tau_P \sim c_{\text{NN}}^{-1}, \]  

evidenced by the lines in Fig. 4. Importantly, the linear scaling of \( \tau_X \) and \( \tau_P \) with \( c_{\text{NN}}^{-1} \) is independent of the dimensionality of the system, which only couples to \( c_B \); see the Appendix for details. Note that the scaling of \( \tau_X \) and \( \tau_P \) with temperature, \( T \), can be more complex due to the coupling between the bulk and nearest-neighbor sites, and this scaling also depends on the dimensionality of the system (Appendix).

The data in Fig. 4 strongly suggests that the exchange and persistence times scale linearly with \( c_{\text{NN}}^{-1} = 1 + \exp(J'/k_B T) \). The energetic coupling \( J' \) captures the net effect of solute-solvent interactions in the solvation shell, such that the water exchange time scales exponentially with the solute-solvent interaction energy. Indeed, Lee, Thirumalai, and Hyeon have recently shown that water exchange times scale exponentially with solute-solvent interactions within the coordination shell, in addition to an entropic contribution, in agreement with our predictions \( \text{[9]} \). The energy scale \( J' \) is model-dependent, and in classical models typically arises from LJ and Coulomb interactions, the latter of which ultimately depends on the charge density of the ion.

The scaling with \( J' \) is simplified under conditions where only the charge density of the ion is changing, with remaining interaction potentials held fixed. In this case, the difference in the energetic coupling between ions arises from electrostatic interactions alone. Such a situation arises for trivalent lanthanides, for example, where the major impact of filling the 4f orbitals is an increase in the strength of solute-solvent electrostatic interactions due to an increase in the charge density (decrease in size) of the ion \( \text{[5]} \). A Born theory-like model for a hydrated ion would suggest that the electrostatic energy contribution to \( J' \) behaves as \( J' \propto Q^2/R \), where \( Q \) and \( R \) are the charge and radius of the ionic solute, respectively \( \text{[22][23]} \). Figure 5 shows that the experimentally-determined water exchange times around trivalent lanthanides follow the expected exponential scaling, \( \ln \tau \sim J'/R \). Note that we have limited our analysis to ions for which well-defined exchange times have been determined \( \text{[5]} \).

Our results are consistent with the system being in close proximity to a phase transition in trajectory space \( \text{[19][10][55]} \). This is supported through the examina-
We have demonstrated that water exchange reactions exhibit facilitated dynamics and may be described within the framework of dynamic facilitation theory. Through the development of a kinetically constrained model, we predict that water exchange times — the inverse of the exchange rate — scale exponentially with the strength of solute-water interactions, consistent with the ultrasensitivity of water exchange times observed experimentally and recent empirical findings. We conclude by noting that the kinetically constrained models discussed here can be readily extended to study solvent exchange in glassy matrices by imposing kinetic constraints in both the solvent and solute nearest-neighbor regions, with different effective temperatures in the two regions. Moreover, extensions to two- and three-dimensional lattices will enable the investigation of the phenomenology underlying directional solvent exchange and solvent exchange at interfaces, which are of importance in geochemical processes and electrochemical processes, for example, as well as solvent transport through porous solids. Further extension to multi-site lattice models may also facilitate the study of exchange dynamics involving ionic solutes and polymeric solvents. We hope that the connections made here between water exchange reactions, kinetically constrained models, and DF theory will further the development of predictive theories for solvent exchange.

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**Appendix A: Appendix: Scaling of Exchange Times with Excitation Concentration**

Following previous work, we obtain a mean-field estimate for the behavior of $\tau_X$ by noting that the facilitation function for one of the solute nearest neighbor sites is given by $f_i = n_{i+1}$; the other is $f_i = n_{i-1}$. This is the same facilitation function as the 1D East model, and we focus on the first neighbor for simplicity.

We start by noting that the rate for the exchange $0 \to 1$ is given by $k^{(+)} = e^{-1/T^*} f_i$, and the rate for the opposite $1 \to 0$ process is $k^{(-)} = f_i$. The mean exchange time for the $n_i = 0$ state is then

$$\langle t_X(0) \rangle \approx \frac{1}{\langle k^{(+)} \rangle} = \frac{e^{1/T^*}}{c_B},$$

and that for the $n_i = 1$ state is given by

$$\langle t_X(1) \rangle \approx \frac{1}{\langle k^{(-)} \rangle} = \frac{1}{c_B},$$

where we have noted that $\langle f_i \rangle = c_B$. The total mean exchange time is then given by

$$\tau_X \approx \langle t_X \rangle \approx \frac{1}{2} \left[ \langle t_X(0) \rangle + \langle t_X(1) \rangle \right]$$

$$= \frac{1}{2c_N c_B},$$

where we have used $1 + e^{1/T^*} = 1/c_N$. For a general $d$-dimensional square lattice, the average facilitation function is given by

$$\langle f_i \rangle = 1 - (1 - c_B)^{ad},$$

where the product $ad$ indicates the number of sites that can facilitate a change in state. For the bulk FA and East models, $a = 2$ and $a = 1$, respectively, independent of $d$. However, due to the presence of the solute site, $a = a(d)$ is a function of $d$ in our model for solvent exchange. For example, $a(1) = 1$, $a(2) = 2/3$, and $a(3) = 5/3$ when using the kinetic constraints of the FA model. We can then carry out the above derivation using Eq. A5 to yield

$$\tau_X \approx \frac{1}{2c_N c_B [1 - (1 - c_B)^{ad}]},$$

which illustrates that $\tau_X \sim c_N^{-1}$ for all $d$, such that the scaling with solute-solvent interactions is also independent of $d$. Note that the temperature dependence of $\tau_X$ is more complex because both $c_N$ and $c_B$ depend on $T$.

The mean persistence time is related to the moments of the exchange time through

$$\tau_p = \frac{\langle t_X^2 \rangle}{2 \langle t_X^1 \rangle},$$

Using the variance, this can be rewritten as

$$\tau_p = \frac{\langle t_X^2 \rangle}{2 \langle t_X \rangle} \left[ 1 - \frac{\langle t_X^2 \rangle}{\langle t_X \rangle^2} \right] \sim c_N^{-1},$$

where $\delta t_X \equiv t_X - \langle t_X \rangle$.

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