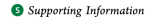


Fragment-Based Quantum Mechanical/Molecular Mechanical Simulations of Thermodynamic and Kinetic Process of the Ru²⁺-Ru³⁺ Self-Exchange Electron Transfer

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ABSTRACT: A fragment-based fractional number of electrons (FNE) approach is developed to study entire electron transfer (ET) processes from the electron donor region to the acceptor region in the condensed phase. Both regions are described by the density-fragment interaction (DFI) method, while FNE as an efficient ET order parameter is applied to simulate the electron transfer process. In association with the QM/MM energy expression, the DFI-FNE method is demonstrated to describe ET processes robustly with the Ru²⁺-Ru³⁺ self-exchange ET as a proof-of-concept example. This method allows for systematic calculations of redox free energies, reorganization energies, electronic couplings and the absolute ET rate constants within the Marcus regime.

1. INTRODUCTION

Electron transfer (ET) is one of the most fundamental processes in chemistry, biochemistry, and material science. The Marcus theory for electron transfer characterizes the ET rate constant as²

$$k_{\rm ET} = \frac{2\pi}{\hbar} \frac{1}{\sqrt{4\pi\lambda k_{\rm B}T}} \langle H_{\rm DA}^2 \rangle \exp\left(-\frac{(\Delta G + \lambda)^2}{4\lambda k_{\rm B}T}\right) \tag{1}$$

where H_{DA} is the donor-acceptor electronic coupling, the symbol $\langle ... \rangle$ denotes the thermal average, λ is the reorganization energy, ΔG is the ET reaction free energy, $k_{\rm B}$ is Boltzmann's constant, and T is the temperature (298 K in this work). Although the Marcus theory is semiclassical and is based on transition-state theory, which assumes weak coupling and slow solvation limit,³ it has been applied successfully to elucidate and compute the ET rate constants for many chemical systems.⁴⁻⁷ As such, accurate calculations of each term in eq 1 are pivotal to facilitating the understanding of the ET process with atomistic and electronic details. To calculate the parameters, e.g. $\langle H_{\rm DA}^2 \rangle$, ΔG , and λ , for ET processes, many methods have been developed using empirical, ^{8,9} semiclassical, or quantum mechanical models ^{10–18} and classical ^{19–24} or density functional based MD simulations. ^{25–47} A more detailed discussion of these works is beyond the scope of this work but has been summarized in a number of reviews. ^{48–55} We note that only few practical *ab initio* methods, ^{37,43} using constrained density functional theory (DFT)⁵⁶ or time dependent DFT, have been proposed to systematically compute the key terms: $\langle H_{\mathrm{DA}}^{2} \rangle$, ΔG , λ , and the final ET rate constant. Usually, these terms are calculated using different schemes. For instance, to obtain the ensemble average of H_{DA}^{2} , classical molecular mechanical (MM) molecular dynamics (MD) simulations are frequently performed to explore the conformational ensemble of either the product or reactant state, without sampling the adiabatic ET transition state. For ΔG and λ , methods using a combination of

QM treatments and continuum models have been commonly employed to approximate the explicit environments, but they inevitably miss some details of the environments, including entropic, dynamic, and conformational contributions, which are especially important for protein systems. 18,27 In order to reach a theoretical estimation of $k_{\rm ET}$, one must combine various schemes, and ensure each scheme can yield results directly comparable to experiments. Therefore, developing a systematic method is still important for understanding the microscopic kinetics, thermodynamics, and mechanism of ET processes at the atomistic and electronic level.

A challenging task for theoretical simulations on k_{ET} is how to balance the computational cost and accuracy. The hybrid quantum mechanical and molecular mechanical (QM/MM) method becomes a very promising tool to simulate the ET process: QM describes the region involving electron transfer while the rest of the system is simulated by MM. In our previous work, we demonstrated that, using the fractional number of electrons (FNE) as the order parameter in the "onthe-fly" QM/MM simulations, 57 the redox potential residing in ΔG could be computed accurately and the ET half reaction processes could be characterized as well. When combined with the QM/MM minimum free energy path (MFEP) method, 58,59 the resulting QM/MM-MFEP/FNE can simulate the ET half reactions for organic molecules and proteins because of the efficiency gained. 60 However, the FNE approach has been limited to the study of the ET half reaction rather than the entire ET process from the electron donor region to the acceptor region. As such, the electronic coupling term or reorganization energy term of ET cannot be computed readily using FNE.

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In this work, we combine the density-fragment interaction (DFI) method with FNE (i.e., DFI-FNE) to simulate the entire ET transfer process in association with "on-the-fly" QM/MM simulations. The DFI method, based on the Kohn-Sham density functional theory (KS-DFT), was originally developed to achieve the linear scaling of computational cost with respect to the growing system size. 61 In particular, we reformulated DFI to reduce the computational cost and make it applicable for large molecular systems.⁶² For instance, we demonstrated that our new DFI method can simulate the thermodynamic properties of liquid water efficiently and accurately. DFI is essentially the simplest divide-and-conquer (DAC)^{63,64} method without any buffer region, but with frozen fragment pseudopotentials. In DFI, the entire QM system is first divided into small fragments. Then each fragment is computed independently, and the desired properties of the entire system are evaluated using the solved fragment information. Interestingly, the fragmentation process can be viewed as the electron localization step. For each fragment in DFI, the number of electrons is fixed and localized in its own region. Therefore, we can gradually move one electron from one fragment to another fragment using FNE as the order parameter to simulate the entire ET process while the environment can be simulated by MM.

This paper is organized as follows. We first briefly review the DFI and FNE approaches and introduce the combined DFI-FNE QM/MM scheme for ET simulations. In section 3, we show the computational details of our simulations on the $\mathrm{Ru^{2+}}\mathrm{-Ru^{3+}}$ self-exchange ET process in aqueous solution as a simple but essential testing case. In section 4, we elucidate how to compute $\langle H_{\mathrm{DA}}^2 \rangle$, ΔG , and λ to obtain k_{ET} and discuss some challenging issues related to the accuracy. Finally, we conclude our work in section 5.

2. METHODS

2.1. Density Fragment Interaction (DFI) Approach. Briefly, we reformulated the DFI energy expression as

$$E[\rho] = \sum_{i=1}^{N_{\rm F}} E_i^F[\rho_i] + E_{\rm ESP}^{\rm FF}[\{\rho_i\}]$$
 (2)

Here, $\rho(\mathbf{r}) = \sum_{i=1}^{N_F} \rho_i(\mathbf{r})$ and ρ_i is the electron density of fragment *i*. $E_i^F[\rho_i]$ is the self-energy of fragment *i*

$$E_{i}^{F}[\rho_{i}] = T_{s}[\rho_{i}] + \int V_{i}(\mathbf{r}) \rho_{i}(\mathbf{r}) d\mathbf{r} - \int \sum_{j \neq i}^{N_{F}} V_{j}^{ESP}(\mathbf{r}) \rho_{i}(\mathbf{r})$$
$$d\mathbf{r} + \int \sum_{j \neq i}^{N_{F}} V_{j}^{FFP}(\mathbf{r}) \rho_{i}(\mathbf{r}) d\mathbf{r} + J[\rho_{i}] + E_{xc}[\rho_{i}]$$
$$+ E_{NN}^{i}$$
(3)

 $E_{\mathrm{ESP}}^{\mathrm{FF}}[\{\rho_i\}]$ is the electrostatic interaction energy between fragments:

$$E_{\text{ESP}}^{\text{FF}}[\{\rho_i\}] = \frac{1}{2} \sum_{N_F}^{i=1} \int \rho_i(\mathbf{r}) \sum_{N_F}^{j \neq i} V_j^{\text{ESP}}(\mathbf{r}) d\mathbf{r}$$
$$+ \frac{1}{2} \sum_{N_F}^{i=1} \sum_{n_i}^{a=1} Z_a^i \sum_{N_F}^{j \neq i} V_j^{\text{ESP}}(\mathbf{R}_a^i)$$
(4)

 $N_{\rm F}$ is the number of fragments in the system, and n_i is the number of atoms in fragment *i*. Z_a^i is the nucleic charge of atom

a in fragment i, and \mathbf{R}_a^i is the corresponding nucleic position. In our new DFI reformulations, we introduced $V_j^{\text{FFP}}(\mathbf{r})$, the frozen fragment pseudopotential for atom j, which takes the Pauli repulsive interactions among fragments into account and approximated $V_j^{\text{ESP}}(\mathbf{R}_a^i)$ using the simple ESP charges. Note that the frozen fragment pseudopotential depends on atomic type and needs to be fitted using a set of training molecules. In this work, we fitted the Pauli parameters for Ru, O, and H with BLYP/Lanl2dz using the fitting protocol described in ref 62. The final parameters used in this work are listed in Supporting Information (SI) Table S1. Note that these parameters are negligible when the edge to edge distance between donor and acceptor groups is beyond 5 Å.

2.2. Fractional Number of Electrons (FNE) Approach. The FNE approach is illustrated in Figure 1 for removing one

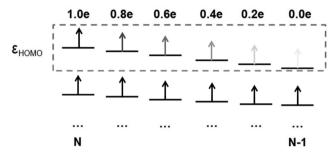


Figure 1. Schematic view of FNE procedure.

electron from the electron donor group to simulate the ET half reaction

$$A_{aq} \rightarrow A_{aq}^+ + e^- \tag{5}$$

Here, FNE is the order parameter to drive the system from one state to the other to achieve the sufficient sampling of free energy. In contrast to mixing two potential energy functions to describe the transition from one state to the other, 28,29,31,33 FNE gradually changes the number of electrons in the system, which only requires one QM calculation for a given conformation. Figure 1 shows that the number of electrons on the HOMO denoted by η is gradually decreased from 1.0 to 0.0. According to the Janak theorem and its extension to generalized Kohn–Sham calculations in our recent work, 57,66 the energy derivative with respect to the number of electrons (expressed as the frontier occupation number) is

$$\frac{\partial E}{\partial \eta} = \varepsilon_{\text{HOMO}} \tag{6}$$

for both Kohn–Sham calculations with explicit density functionals and generalized Kohn–Sham calculations with an orbital functional. Thus, the free energy change (i.e., oxidation free energy) of eq 5 can be readily computed by thermodynamic integration (TI):

$$\Delta A = -\int_0^1 \frac{\partial A(\eta)}{\partial \eta} d\eta = -\int_0^1 \langle \frac{\partial E(\eta)}{\partial \eta} \rangle_{\eta} d\eta$$
$$= -\int_0^1 \langle \varepsilon_{\text{HOMO}} \rangle_{\eta} d\eta$$
(7)

The FNE-TI approach has been applied successfully to compute the redox potential of the metal ion in aqueous solution. In addition, the reorganization energy of the half reaction and activation free energy of the full reaction

(assuming two reaction centers do not interact with each other) can be calculated by reusing the sampled trajectories using the energy gap ΔE as the reaction coordinate.⁵⁷

2.3. Combined DFI-FNE QM/MM Simulations for Full ET Reactions. To simulate the realistic ET reaction process, the previous half reactions studied by FNE are not satisfactory since the electron must be driven from the donor to the acceptor to characterize the ET mechanism with both donor and acceptor groups present. Here, we combine the DFI method with FNE and TI to simulate the full ET process. In this work, we focus on using a simple ET reaction of Ru²⁺–Ru³⁺ self-exchange in aqueous solution to demonstrate how DFI-FNE works:

$$Ru_{D}^{2+}(H_{2}O)_{6} + Ru_{A}^{3+}(H_{2}O)_{6}$$

$$\rightarrow Ru_{D}^{3+}(H_{2}O)_{6} + Ru_{A}^{2+}(H_{2}O)_{6}$$
(8)

As shown in Figure 2, for a given distance between two ruthenium metal centers, each hydrated Ru is one independent

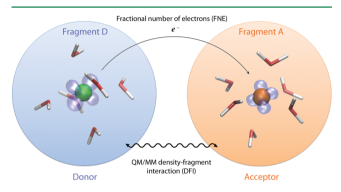


Figure 2. Schematic view of DFI-FNE procedure for the $\mathrm{Ru}^{2+}\mathrm{-Ru}^{3+}$ self-exchange ET process.

fragment in DFI. As such, it is natural to use FNE as the order parameter to drive the electron from one Ru to the other since the number of electrons in each fragment can be specified (i.e., the electrons are naturally localized within fragments). Therefore, the free energy change of the full ET reaction such as eq 7 can be obtained by TI:

$$\Delta A = -\int_0^1 \frac{\partial A(\eta)}{\partial \eta} d\eta = -\int_0^1 \left\langle \frac{\partial E^{\text{DFI}}(\eta)}{\partial \eta} \right\rangle_{\eta} d\eta$$
$$= -\int_0^1 \left\langle \varepsilon_{\text{HOMO}}^{\text{D}} - \varepsilon_{\text{HOMO}}^{\text{A}} \right\rangle_{\eta} d\eta \tag{9}$$

where $\varepsilon_{\text{HOMO}}^{\text{D}}$ is the HOMO energy of fragment D and $\varepsilon_{\text{HOMO}}^{\text{A}}$ is the HOMO energy of fragment A. Note that the HOMO becomes LUMO in fragment D at the finial state while the LUMO becomes HOMO in fragment A after the fractional electron is moved from fragment D to fragment A. ΔA vanishes in the specific self-exchange ET, but finite free energy differences are also allowed in this method for general cases.

By analyzing the sampled DFI-FNE trajectories, we can compute $\langle H_{\mathrm{DA}}^{2} \rangle$ and λ required in eq 1 using the consistent DFI QM/MM energy expression without introducing any *ad hoc* parameter. The reorganization energy λ can be readily evaluated by using the energy gap as the reaction coordinate. ¹⁹ For H_{DA} , we apply the fragment orbital approach combined with DFI. The frontier molecular orbitals for each fragment can

be extracted and taken as the donor and acceptor orbital (i.e., $\varphi_{\rm D}$ and $\varphi_{\rm A}$) to compute $H_{\rm DA}$: ⁶⁷

$$H_{\rm DA} = T_{\rm DA} - S_{\rm DA} (T_{\rm DD} + T_{\rm AA})/2$$
 (10)

where $T_{\rm DA} = \langle \varphi_{\rm D} | F \hat{\varphi}_{\rm A} \rangle$, $T_{\rm DD} = \langle \varphi_{\rm D} | F \hat{\varphi}_{\rm D} \rangle$, $T_{\rm AA} = \langle \varphi_{\rm A} | F \hat{\varphi}_{\rm A} \rangle$, and $S_{\rm DA} = \langle \varphi_{\rm D} | \varphi_{\rm A} \rangle$. The operator \hat{F} is the Fock operator that is constructed from the density of the entire system ρ without fragmentations. Total density ρ is the summation of converged fragmental densities $\rho_{\rm D}$ and $\rho_{\rm A}$ obtained from self-consistent DFI calculations. Note that our coupling calculations are similar to the FO-DFT used in a previous work. However, the DFI frontier molecular orbitals are used here, rather than the CDFT wave functions, 39,40 to represent the two diabatic states for the donor and acceptor, respectively. In order to obtain $H_{\rm DA}$ with consistent relative phases between the donor and acceptor orbital, we adjust the signs of the donor and acceptor orbital, respectively, in each MD snapshot to be the same as the signs in the first snapshot.

The Ru^{2+} – $\hat{R}u^{3+}$ self-exchange ET process is simple but extremely challenging. Normal QM computations with both ions are hardly even converged due to electronic state degeneracy and delocalization errors of functionals. Moreover, the ET rate constant of the Ru^{2+} – Ru^{3+} self-exchange process requires extensive sampling with different Ru^{2+} – Ru^{3+} distances. Hence, the final rate constant needs to be computed and integrated as I^{7+}

$$k_{\text{total}} = \int_0^\infty 4\pi r^2 e^{-G(r)/k_B T} k_{\text{ET}}(r) dr$$
 (11)

where G(r) is the potential of mean force for the two reactant complexes in solution that determines the pair distribution of the donor and acceptor. We will elucidate how DFI-FNE can be applied in this case and what are challenges in DFI-FNE.

3. COMPUTATIONAL DETAILS

The self-exchange redox couple $Ru(H_2O)_6^{2+}-Ru(H_2O)_6^{3+}$ was simulated with the DFI method with two individual fragments. The initial geometries of the two Ru complexes were built 10.0 Å apart, then solvated using TIP3P explicit solvent model in a cubic water box with the side length of 64 Å. A harmonic restraint was applied on the two Ru atoms with the equilibrium distance of 10.0 Å and a force constant of 100 kcal/mol/Å². The separation distances were then sequentially reduced to 6.0 Å with an increment of 1.0 Å in a series of MD simulations that generated the initial geometries of Ru2+-Ru3+ at different distances. Since the radii of the aqueous Ru^{2+/3+} complexes are determined to be 5.5-6 Å, a system with a separation distance of less than 6 Å would cause significant electronic coupling between the donor and acceptor, undermining the decoupled assumption in DFI. Thus, DFI calculations with a separation distance of less than 6 Å is avoided. The system was equilibrated for 16 ps with a timestep of 1 fs using the NVT ensemble, with a reduced restraining force constant of 10 kcal/ mol/Å². In the following production MD, the simulation timescale is 16 ps for each separation distance, with 1000 snapshots recorded to compute the H_{DA} and reorganization energies. The DFI method with BLYP/LanL2DZ was used for the QM subsystem in the production MD, and BLYP/ LanL2DZ(Ru)+6-31+G*(H₂O) was used in the calculation of H_{DA} . Classical MM MD umbrella samplings with force field parameters adapted from ref 34 were performed to obtain the potential mean force (PMF) of the Ru-Ru distance with a force cutoff of 16 Å combined with the particle mesh Ewald

method, since the long-range electrostatic interactions caused by periodic boundary conditions (PBC) have not been incorporated in our QM/MM simulations. These long-range interactions are important in computing the PMF profile for two ions. ^{44,72,73} In the PMF umbrella sampling, a restraining potential of 5 kcal/mol/Ų was used to increase the overlap between sampling windows, generating an average of standard deviations of 0.35 Å for the Ru–Ru distance. The window interval of 0.5 Å was used to cover the distances ranging from 5.5 Å to 16 Å, and 160 ps MD was carried out for each window. The weighted histogram analysis method was used to reconstruct the PMF. ⁷⁴

4. RESULTS AND DISCUSSION

4.1. Oxidation Free Energy. The absolute oxidation free energies were obtained by integrating the ensemble averages of eigenvalues of the HOMO $\langle \varepsilon_{\text{HOMO}} \rangle$ of the Ru complexes according to eq 6. Figure 3 shows a profile of $\langle \varepsilon_{\text{HOMO}} \rangle$ at

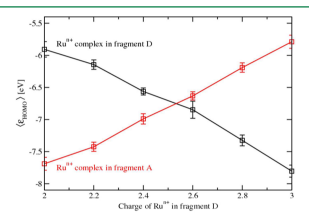


Figure 3. Symmetrical redox profile of the self-exchange ET reaction (Ru–Ru distance of 7.0 Å is shown). Values of $\langle \varepsilon_{\rm HOMO} \rangle$ are obtained in DFI simulations, and the error bars show the statistic uncertainties. Integration of each $\langle \varepsilon_{\rm HOMO} \rangle$ curve generates the oxidation free energy of the corresponding Ru complex.

different FNEs of the Ru-Ru couple at a separation distance of 7.0 Å. The self-exchange ET reaction shows a symmetric profile of the two $\langle \varepsilon_{\mathrm{HOMO}} \rangle$ series. The integrations of the two curves generate the absolute oxidation free energies for the two fragments of 6.75 and 6.79 eV, with a difference of 0.04 eV. The average discrepancy of the oxidation free energy between the two self-exchange fragments is 0.03 eV over various separation distances, which is entirely due to the convergence in the statistical sampling. One may note that the absolute oxidation free energies of the aqueous Ru complex obtained in the full ET scheme are remarkably higher than that calculated in the half ET reaction (4.96 eV).⁵⁷ The shift of the oxidation free energies is mainly caused by the electrostatic field exerted by the second Ru complex in proximity. The presence of the two positively charged Ru complexes raises the electrostatic potential on each other, thus making them stronger oxidizers. As shown in Figure 4, we plot the oxidation free energies as a function of separation distance. The oxidation free energies decay from 7.05 eV at r = 6.0 Å to 6.50 eV at r = 10.0 Å, which fit well to the screened Coulomb potential shown as a dashed line. Note that when long-range electrostatic interactions are considered, the oxidation free energies may be lower.

4.2. Reorganization Energy. The reorganization is a key factor that determines the rate constant and the mechanism of

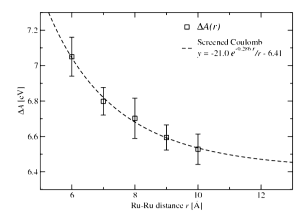


Figure 4. Distance dependence of oxidation free energies of Ru complexes. The calculated results are shown in black squares with error bars showing the sampling uncertainties, and the dashed line is fitted to the screened Coulomb potential.

an ET process. On the basis of the DFI method, we can obtain the reorganization energy of the full ET process by projecting the diabatic free energy profiles over the energy gap similarly to the procedure in a half ET reaction. ⁵⁷ The essential difference is that the energy gap of the full ET reaction is the energy difference between the swapped electronic states of the redox couple, i.e.

$$\Delta E = E[D(Ru^{2+}) - A(Ru^{3+})] - E[D(Ru^{3+}) - A(Ru^{2+})]$$
(12)

Because the QM/MM electrostatic interaction cutoff is used in the DFI simulations, we investigated the influence of the choice of different cutoff radii on the final values of the reorganization energies. We chose two extreme separation distances, 6.0 Å and 10.0 Å, to represent the two scenarios of close proximity and long ET distance. For each separation distance, cutoff distances of 8.0, 12.0, and 20.0 Å were examined, and the corresponding reorganization energies are shown in Figure 5. As we can see, the choice of the cutoff distance does not affect the reorganization energy of the Ru–Ru couple at close proximity, while the electrostatic cutoff shows a significant impact on the Ru–Ru couple at long distances. The increased separation distance between the two charged Ru complexes leads to disturbed solvent distributions

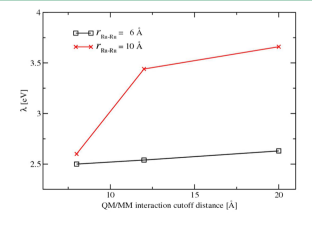


Figure 5. Influence of QM/MM electrostatic cutoff distance on reorganization energies.

in a larger region and thus requires a larger QM/MM interaction cutoff. Furthermore, a comparison of the reorganization energies of the five various separation distances between the choice of a 12.0 Å or 20.0 Å cutoff are shown in Figure S2 and summarized in Table S2. Reorganization energies obtained using a cutoff of 20.0 Å generally show about a 0.1 eV discrepancy compared to the results obtained using 12.0 Å. The λ convergence at a 20.0 Å cutoff is also in agreement with our previous work on half ET reactions where detailed convergence tests were performed.⁵⁷ Therefore, the cutoff distance is set to 20.0 Å in the production runs of DFI simulations.

The individual diabatic free energy profiles showing the reorganization energies at difference separation distances are plotted in Figure S1, and the values of the reorganization energies are summarized in Table 1. In addition to the total

Table 1. Reorganization Energies, H_{DA} and k_{ET} at Different Ru-Ru Distance

r/Å	$\langle H_{\mathrm{DA}}^{2} \rangle^{1/2} / \mathrm{eV}$	$\lambda(r_{\text{cutoff}} = 20 \text{ Å})/\text{eV}$	$k_{\rm ET}/{ m M}^{-1}{ m s}^{-1}$	$\begin{array}{c} \lambda_{scaled}/\\ eV \end{array}$	$k_{\mathrm{ET}} \frac{(\lambda_{\mathrm{scaled}})}{\mathrm{s}^{-1}a}$
6.0	2.03×10^{-3}	2.63	3.81×10^{-1}	1.66	5.88×10^{3}
7.0	3.33×10^{-4}	2.97	3.59×10^{-4}	1.87	1.87×10^{1}
8.0	1.07×10^{-4}	3.23	2.89×10^{-6}	2.03	3.80×10^{-1}
9.0	3.75×10^{-5}	3.40	6.71×10^{-8}	2.14	1.62×10^{-2}
10.0	1.66×10^{-5}	3.66	1.02×10^{-9}	2.31	6.28×10^{-4}

"The free energy difference of the self-exchange process (ΔA) is rounded to zero in the calculations of $k_{\rm ET}$.

reorganization energies, λ can also be decomposed to inner and outer shell reorganization energies by re-evaluating the sampled energies in the absence of solvent molecules.⁵⁷ As an example of the Ru couple at r=8.0 Å, the inner shell diabatic ET surfaces were constructed as shown in Figure S3, which generates the inner shell reorganization energy of 0.69 eV. On the basis of the total reorganization energy of 3.13 eV, we can derive that the outer shell contribution is 2.44 eV, which is a major part of the reorganization energy in the Ru²⁺–Ru³⁺ self-exchange ET reaction. This suggests that the outer shell reorganization plays important roles in determining the total reorganization energy since two hydrated ions are rigid during ET.

We noticed that the reorganization energies calculated using our QM/MM method are consistently overestimated. In the comparison of other simulations on the same self-exchange process, we find that the lack of a polarizable solvent model is an important reason for the overestimation. In addition, the simulations with solvent polarization generate relatively lower and more accurate reorganization energies. And addition, the lack of PBC in QM calculations may also contribute to the error in the reorganization energies. Since the reorganization energy has an exponential dependence on ET rate $k_{\rm ET}$ (for self-exchange reactions with zero redox free energy), we here rescaled the reorganization energy to reduce the errors due to the lack of solvent polarization and PBC with the scaling coefficient 0.63, which was determined from the ratio of the reorganization energy of a single Ru complex (2.45 eV) from our previous work to the experimental estimation (1.54 eV).

4.3. Electronic Coupling. To calculate the ET rate, the electronic coupling $H_{\rm DA}$ needs to be determined. According to our DFI scheme, we computed the electronic coupling using the fragment orbital approximation (eq 10). Since the

calculation of $H_{\rm DA}$ involves the molecular orbital overlap $S_{\rm DA}$, we examined the basis set dependence of $S_{\rm DA}$. The basis set LanL2DZ is used in the direct MD sampling to achieve affordable computational cost for the "on-the-fly" DFI-FNE/MM calculations. On the basis of the MD trajectories, we reevaluated the $S_{\rm DA}$ of each snapshot using larger basis sets, 6-31+G* and 6-31++G**. The comparison of $|S_{\rm DA}|$ among different basis sets is shown in Figure 6. The other two large

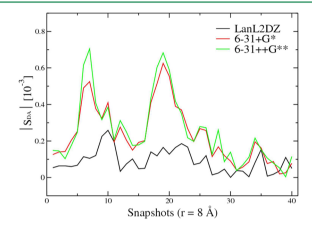


Figure 6. Basis set dependence of $|S_{DA}|$ using LanL2DZ, 6-31+G*, and 6-31++G**. Convergence is achieved at 6-31+G*.

basis sets show good agreement with each other, indicating the convergence of the size of the basis set; however, the small basis set LanL2DZ is obviously not sufficient to calculate accurately the orbital overlap S_{DA} and the electronic coupling H_{DA} . As such, we choose the 6-31+G* in the calculations of $H_{\rm DA}$ to balance the computational cost and the accuracy. In addition, we analyzed S_{DA} in different ET reaction regions by comparing the ensemble averages of S_{DA}^2 obtained from samplings at different electronic states as shown in Figure 7. On the basis of the conformational trajectories of reactant and product states, the values of $\langle S_{DA}^2 \rangle$ at the reaction end points are about 10^{-9} , which is 1 order of magnitude smaller than the ensemble averages obtained for the diabatic transition state ($\sim 10^{-8}$) where ΔE < 0.05 eV. Although the FNE state with an even distribution of electrons on the donor and acceptor (Ru^{2.5+}-Ru^{2.5+}) restrains the system near the transition state ($\langle \Delta E \rangle$ = 0.002 eV), it still allows thermal fluctuations of the solvent dynamics and shows the standard deviation of ΔE to be 0.37 eV. Therefore, the ensemble of Ru^{2.5+}-Ru^{2.5+} is not the strict diabatic transition state ensemble (defined as $|\Delta E|$ < 0.05 eV). Interestingly, as shown in Figure 7, the statistics of $\langle S_{DA}^2 \rangle$ are almost identical for the two ensembles, indicating that the FNE state is a good approximation to the diabatic ET transition

Strictly, the electronic coupling $H_{\rm DA}$ between the donor and acceptor fragments should also be computed at the transition states, the cross region where the energy gap is zero. Thus, only the snapshots with energy gaps less than 0.05 eV were used to compute the ensemble averages of $H_{\rm DA}$. The final values of $H_{\rm DA}$ for the Ru couple at different separation distances were summarized in Table 1. The distance dependence of $H_{\rm DA}$ was analyzed and fitted to exponential distance dependence $|H_{\rm DA}| = A \exp[-\beta/2]$, with the distance decay constant $\beta/2 = 1.16~{\rm \AA}^{-1}$, as shown in Figure 8. Previous simulations on the same ET process reported similar values and a distance dependent decay rate of $H_{\rm DA}$. The distance decay rate constant in our

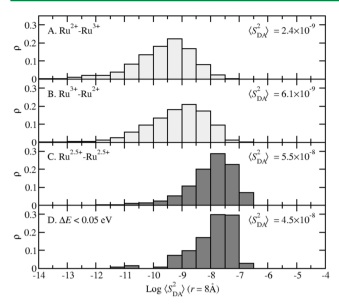


Figure 7. Distribution of $\langle S_{\mathrm{DA}}^2 \rangle$ in various electronic states: (A) reactant state (light gray), (B) product state (light gray), (C) adiabatic FNE state with even electron distribution (dark gray), (D) diabatic transition state with energy gap criterion of $\Delta E < 0.05$ eV (dark gray). The ensemble average value of $\langle S_{\mathrm{DA}}^2 \rangle$ shows about a 1 order of magnitude increase in the transition state compared to the reactant or product state.

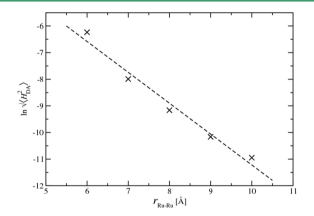


Figure 8. Distance dependence of $H_{\rm DA}$ (black cross). Linear regression (dashed-line) generates the slope of 1.16, which leads to a decay constant β of 2.32 Å⁻¹ (6-31+G* basis set is used to calculate $H_{\rm DA}$.).

simulations agrees with the estimation in aqueous Fe^{2+} – Fe^{3+} self-exchange $(\beta/2=1.2~\text{Å}^{-1})$, and it is significantly larger than those in protein environments $(\beta/2=0.55~\text{Å}^{-1})$. This is not surprising because the Ru^{2+} – Ru^{3+} self-exchange in aqueous solution does not have a specific "bridge" but dynamic water molecules that mediate the ET; hence the ET rate decays much faster when the separation distance increases.

4.4. ET Rate $k_{\rm ET}$ and Total Reaction Rate k of ${\rm Ru}^{2+}$ – ${\rm Ru}^{3+}$ Self-Exchange. With the redox free energy ΔG , the reorganization energy λ , and the electronic coupling $H_{\rm DA}$ obtained individually, we can calculate the ET rate $k_{\rm ET}$ according to eq 1. The values of $k_{\rm ET}$ calculated using both the native and rescaled reorganization energies at different separation distances were summarized in Table 1. The ET rates $k_{\rm ET}$ are greatly promoted using the rescaled reorganization energies. With all the distance dependences, we can finally compute the total reaction rate k by integrating the ET rate $k_{\rm ET}$

over the space following eq 11, where $k_{\rm ET}$ values are averaged with the Ru–Ru pair distribution ${\rm e}^{-G(r)/k_{\rm B}T}$ at the corresponding separation distance. The distance dependence of the intermolecular interactions G(r) sampled by classical MD simulations is shown in Figure 9. The intermolecular

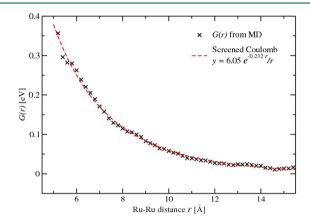


Figure 9. PMF along Ru–Ru distance from classic MD simulations. Simulation results of the Ru–Ru interaction (black cross) behave like a screened Coulomb potential (fitted as red dashed-line).

interaction G(r) agrees with previous simulation results³⁷ and fits the screened Coulomb potential nicely, shown as a dashed line. The baseline of G(r) was offset to zero at $r = \infty$. Spherical integration of the weighted $k_{\rm ET}$ generates the total rate constant of the self-exchange ET reaction between the $Ru^{2+}-Ru^{3+}$ couple as $k=0.11~M^{-1}~s^{-1}$. Before we can directly compare the computed rate constant to experimental measurements, the effect of nuclear tunneling over the activation barrier and the ionic strength must be considered. In the aqueous Fe²⁺-Fe³⁺ self-exchange, the nuclear tunneling introduces a rate enhancement factor of about $10^{2,22,34}$ based on different water models. This increases the total rate constant k for about 1 order of magnitude. In addition, the high ionic concentration in acidic solution may increase the measured ET rate. The overall effect of finite ionic strength (5 M) was reported to promote the rate by about 2 orders of magnitude compared to infinite dilution limit.³⁷ Considering the ~10³ enhancement due to the quantum tunneling and ionic strength, the total rate of 0.11 M^{-1} s⁻¹ in our simulation is scaled to 1×10^2 and shows good agreements with the experimental observation of 20 M⁻¹ s⁻¹.

5. CONCLUSIONS

In this work, we developed the DFI-FNE QM/MM approach to study the full ET reaction. The new method allows us to calculate the redox free energies and reorganization energies of the donor and acceptor simultaneously and generates the redox free energy and diabatic free energy surface of the full ET reaction. Combined with the fragment-orbital approach, the electronic couplings between the donor and acceptor can be readily computed by the DFI scheme. As such, the absolute ET rate constant $k_{\rm ET}$ can be obtained without introducing *ad hoc* parameters using DFI-FNE. This approach should have a significant impact on computing the ET rate constants for chemical and biological systems.

To further improve the accuracy, the long-range electrostatic interactions need to be considered. In addition, how to improve the solvent description is also important in order to obtain accurate reorganization energy. For instance, the MM model in

DFI-FNE QM/MM significantly overestimates λ . More importantly, many other factors in electrochemical experiments, e.g., pH dependence, ion strength, and interfacial effects, can prevent good agreement between theoretical predictions and experimental measurements. The difference between physiological environment and *in vitro* experiments introduce further complications in understandings of many important biochemical ET reactions. Further studies related to these issues will be investigated in our group.

ASSOCIATED CONTENT

S Supporting Information

Figures S1, S2, and S3 and Tables S1, S2, and S3. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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