

## Steering Electrons on Moving Pathways

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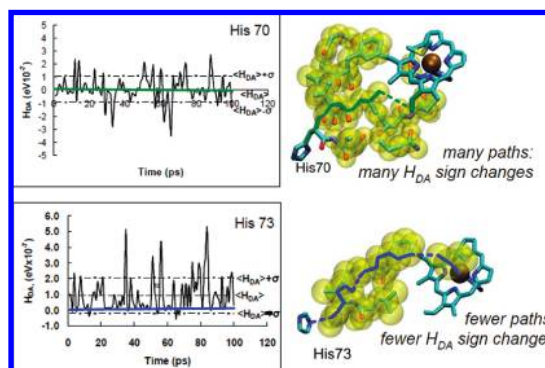
RECEIVED ON APRIL 18, 2009

### CONSPICUOUS

**E**lectron transfer (ET) reactions provide a nexus among chemistry, biochemistry, and physics. These reactions underpin the “power plants” and “power grids” of bioenergetics, and they challenge us to understand how evolution manipulates structure to control ET kinetics. Ball-and-stick models for the machinery of electron transfer, however, fail to capture the rich electronic and nuclear dynamics of ET molecules: these static representations disguise, for example, the range of thermally accessible molecular conformations. The influence of structural fluctuations on electron-transfer kinetics is amplified by the exponential decay of electron tunneling probabilities with distance, as well as the delicate interference among coupling pathways. Fluctuations in the surrounding medium can also switch transport between coherent and incoherent ET mechanisms — and may gate ET so that its kinetics is limited by conformational interconversion times, rather than by the intrinsic ET time scale. Moreover, preparation of a charge-polarized donor state or of a donor state with linear or angular momentum can have profound dynamical and kinetic consequences. In this Account, we establish a vocabulary to describe how the conformational ensemble and the prepared donor state influence ET kinetics in macromolecules. This framework is helping to unravel the richness of functional biological ET pathways, which have evolved within fluctuating macromolecular structures.

The conceptual framework for describing nonadiabatic ET seems disarmingly simple: compute the ensemble-averaged (mean-squared) donor–acceptor (DA) tunneling interaction,  $\langle H_{DA}^2 \rangle$ , and the Franck–Condon weighted density of states,  $\rho_{FC}$ , to describe the rate,  $(2\pi/\hbar)\langle H_{DA}^2 \rangle \rho_{FC}$ . Modern descriptions of the thermally averaged electronic coupling and of the Franck–Condon factor establish a useful predictive framework in biology, chemistry, and nanoscience. Describing the influence of geometric and energetic fluctuations on ET allows us to address a rich array of mechanistic and kinetic puzzles. How strongly is a protein’s fold imprinted on the ET kinetics, and might thermal fluctuations “wash out” signatures of structure? What is the influence of thermal fluctuations on ET kinetics beyond averaging of the tunneling barrier structure? Do electronic coupling mechanisms change as donor and acceptor reposition in a protein, and what are the consequences for the ET kinetics? Do fluctuations access minority species that dominate tunneling? Can energy exchanges between the electron and bridge vibrations generate vibronic signatures that label some of the D-to-A pathways traversed by the electron, thus eliminating unmarked pathways that would otherwise contribute to the DA coupling (as in other “which way” or double-slit experiments)? Might medium fluctuations drive tunneling–hopping mechanistic transitions? How does the donor-state preparation, in particular, its polarization toward the acceptor and its momentum characteristics (which may introduce complex rather than pure real relationships among donor orbital amplitudes), influence the electronic dynamics?

In this Account, we describe our recent studies that address puzzling questions of how conformational distributions, excited-state polarization, and electronic and nuclear dynamical effects influence ET in macromolecules. Indeed, conformational and dynamical effects arise in all transport regimes, including the tunneling, resonant transport, and hopping regimes. Importantly, these effects can induce switching among ET mechanisms.



## Time Scales in Biological Electron Transfer

The golden rule, Born–Oppenheimer, Franck–Condon (FC) formulation of electron transfer (ET) (with rate constant  $k_{\text{ET}} = (2\pi/\hbar)\langle H_{\text{DA}}^2 \rangle \rho_{\text{FC}}$ ) is appropriate when the ET mechanism is coherent electron tunneling from D to A, when the FC times (the amount of time D and A electronic states remain resonant) are shorter than ET coupling fluctuation times,<sup>1–13</sup> when medium relaxation of the product state is fast compared with the ET rate,<sup>14</sup> and when conformational exchange among the “dominant” conformations is rapid on the ET time scale (i.e., the process is not gated).<sup>15</sup> Although we have found that fast bond angle fluctuations (tens of femtoseconds) and slower timescale torsion angle fluctuations of the tunneling bridge have a significant effect on the ET coupling,<sup>13</sup> their contribution to the ET rate for most biological ET reactions is well described by the quantity  $\langle H_{\text{DA}}^2 \rangle$  because such fluctuations are faster than most long-distance ET rate timescales. Other slow events on the time scale of ET may lead to kinetic gating.<sup>16</sup>

Are these conventional approximations in all biological ET systems? In the case of Ru-azurins,<sup>13</sup> decay times of the DA coupling autocorrelation function ( $\langle H_{\text{DA}}(t)H_{\text{DA}}(0) \rangle / \langle H_{\text{DA}}^2(0) \rangle$ ) are tens of femtoseconds, an order of magnitude larger than the time scale for nuclear motion out of the crossing region of the DA potential energy surfaces ( $\sim \hbar/(\lambda k_{\text{B}}T)^{1/2}$ ),<sup>17</sup> where  $\lambda$  is the reorganization energy. For  $\lambda$  on the scale of eV, this time scale is a few femtoseconds, and  $H_{\text{DA}}$  does not change during nuclear wave packet motion through the crossing point of the DA potential energy surfaces. Thus, the ET rate is proportional to  $\langle H_{\text{DA}}^2 \rangle$ ,<sup>12,13,18,19</sup> where the averaging is over the ensemble of protein structures.

The shortest decay time of  $\langle H_{\text{DA}}(t)H_{\text{DA}}(0) \rangle / \langle H_{\text{DA}}^2(0) \rangle$  in azurin was found to arise largely from valence angle fluctuations.<sup>13</sup> Since these valence angle vibrations occur on similar time scales in proteins, the Condon approximation is likely to be valid more broadly for protein ET. However, the validity of the Condon approximation for DNA ET remains an open question.

## The Persistence of Structural Memory in Biological Electron Transfer

The tunneling pathway model of biological ET estimates the DA interaction based on the decay through a sequence of covalent and noncovalent contacts between donor and acceptor. The decay parameters for the contacts along the pathway are based on simple rules derived from the physical principle that bond-mediated interactions are generally larger than through-space interactions.<sup>20</sup> Although simple, this model successfully predicts the dependence of rate on medium structure in dozens of biological and semibiological systems<sup>8,21</sup>

and provides a quick survey of electronic communication for newly reported protein X-ray and NMR structures.

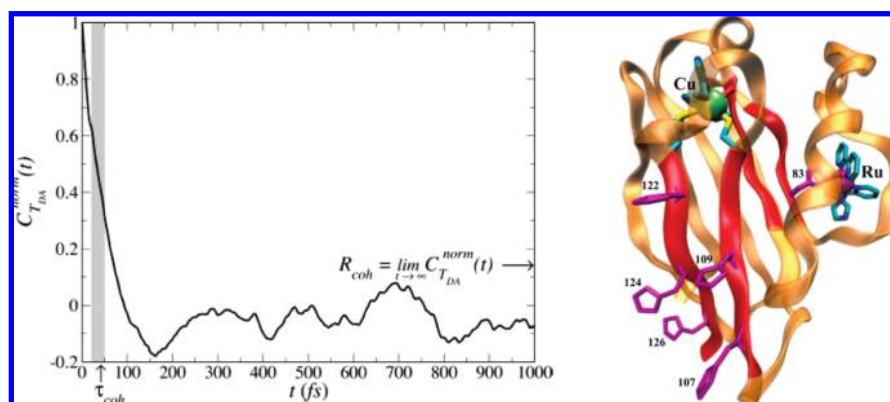
Going beyond the pathway description of electron tunneling mediation, in order to incorporate more specific chemical and multipath effects, especially quantum interferences among paths, necessitates the inclusion of multiple coupling routes and ensemble averaging.<sup>22</sup> Inclusion of quantum interference effects in  $H_{\text{DA}}$  is usually accomplished using effective Hamiltonian and Green’s function methods,<sup>23,24</sup> the generalized Mulliken–Hush scheme,<sup>25,26</sup> and the tunneling current approach.<sup>27</sup>

The mean-squared coupling  $\langle H_{\text{DA}}^2 \rangle$  in  $k_{\text{ET}}$  is the sum of the average coupling squared and the squared coupling variance:

$$\langle H_{\text{DA}}^2 \rangle = \langle H_{\text{DA}} \rangle^2 + \sigma^2 \quad (1)$$

where  $\sigma^2 = \langle (H_{\text{DA}} - \langle H_{\text{DA}} \rangle)^2 \rangle$ . To what extent is the mean-squared coupling dominated by the average coupling compared with fluctuations away from the average? Balabin and Onuchic first posed this question in the context of photosynthetic reactions and examined the relative values of  $\sigma$  vs  $\langle H_{\text{DA}} \rangle$ <sup>28</sup> through the “coherence parameter”  $C = \langle H_{\text{DA}} \rangle^2 / \langle H_{\text{DA}}^2 \rangle = [1 + (\sigma^2 / \langle H_{\text{DA}}^2 \rangle)]^{-1}$  (not to be confused with  $C_{\text{TDA}}^{\text{nom}}(t)$  in Figure 1). When  $\langle H_{\text{DA}} \rangle^2 \gg \sigma^2$ , the average medium structure defines the coupling. However, when  $\langle H_{\text{DA}} \rangle^2 \ll \sigma^2$ , protein geometries that differ from the average protein structure provide coupling interactions that are much larger than the interaction provided by the average structure.

Assessment of mean vs fluctuation-dominated couplings has provided new insights to the field of biological ET in the last 5 years. For example, fluctuation-dominated ET was suggested as a possible means to control charge flow and to introduce novel temperature dependencies to the kinetics.<sup>28</sup> We find that ET reactions to edges of hemes or chlorophylls favor a multipathway mechanism ( $\sigma^2$  dominated), while axial coupling to the coordinated metals in these cofactor rings favors a dominant-pathways ( $\langle H_{\text{DA}} \rangle^2$ ) mechanism.<sup>11</sup> The edge-coupled multipathway mechanism helps to rationalize the near exponential decay as a function of distance for photosynthetic reaction center ET,<sup>29,30</sup> while the axial dominant-pathways mechanism explains the ET kinetics in some semibiological Ru-proteins.<sup>11</sup> The emergence of average medium and few pathway behavior deduced here remains a fascinating topic for research. In the tunneling-limited Ru-modified protein systems of Gray and Winkler,<sup>8</sup> all derivatives with anomalously slow rates for their DA distance are in the large  $C$  few-pathway ( $\langle H_{\text{DA}} \rangle^2 \gg \sigma^2$ ) regime and are axially coupled to the heme donor. “Fast” and “average” rate derivatives (for their distance) fall in the multipath regime ( $\langle H_{\text{DA}} \rangle^2 \ll \sigma^2$ ) with

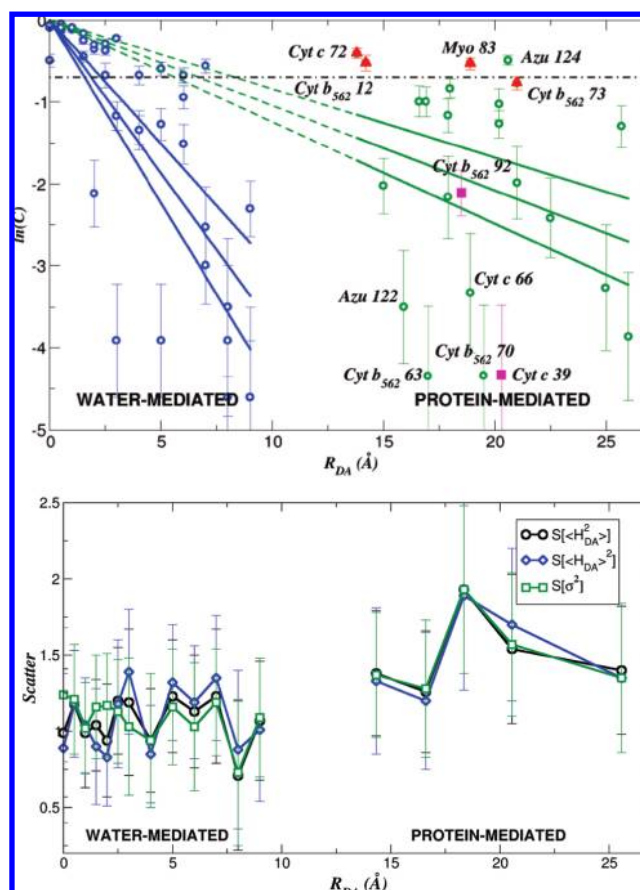


**FIGURE 1.** The normalized electronic coupling autocorrelation function (left)  $C_{T_{DA}}^{norm} = \langle H_{DA}(t)H_{DA}(0) \rangle / \langle H_{DA}^2(0) \rangle$  in Ru-His83 modified azurin decays on the time scale of tens of femtoseconds,<sup>13</sup> 1 order of magnitude longer than the Franck–Condon decay time.<sup>17</sup> During each DA curve crossing event, the DA interaction is essentially unchanged. The fluctuation of  $C_{T_{DA}}^{norm}$ , rapid on the time scale of the ET rate, enter the rate through the mean squared coupling,  $\langle H_{DA}^2 \rangle$ .

effective interactions determined by multiple alternative pathway structures, rather than by a few dominant paths.<sup>11</sup> Examples of  $\sigma^2$ -dominated regimes (with many pathways leading from the heme edge) and  $\langle H_{DA} \rangle^2$ -dominated regimes (with few pathways leading from the heme's axial ligand) appear in ref 11.

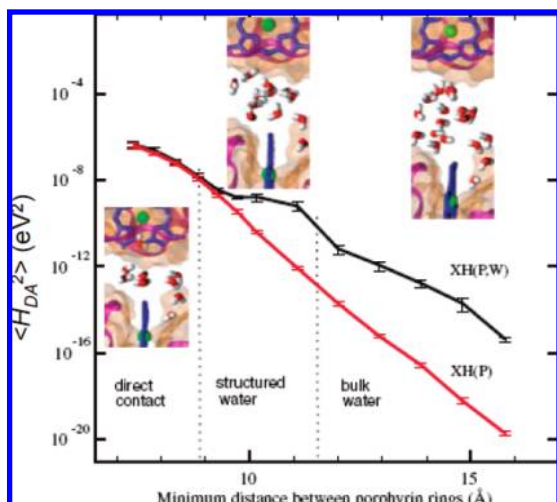
Comparing  $\langle H_{DA} \rangle^2$  to  $\sigma^2$  values in proteins has led us to pose a statistical question regarding coupling: is there a distance that characterizes, on average, a transition between  $\langle H_{DA} \rangle^2$ - and  $\sigma^2$ -dominated kinetics?<sup>31</sup> An answer emerges from coupling analysis for classical molecular dynamics-sampled geometries of Ru-modified proteins,<sup>8</sup> as well as for water-mediated cytochrome *b*<sub>5</sub> self-exchange.<sup>32</sup> We find that, on average, a transition between the  $\langle H_{DA} \rangle^2$  regime and the  $\sigma^2$  regime for protein ET at distances of  $\sim 6$ – $7$  Å, about the size of an amino acid residue (see Figure 2, top). This statistical analysis assumes that the coherence parameter  $C$  is anchored to unity at donor–acceptor contact ( $\ln C = 0$  in Figure 2 at  $R_{DA} = 0$ ). It will be interesting to explore trends in  $C$  at distance under  $6$ – $7$  Å, when tunneling mediation accesses fewer coupling pathways, and also to examine the case of DNA ET. For water-mediated self-exchange, the transition to a fluctuation-dominated coupling regime occurs at distances about the size of a water molecule,  $\sim 2$ – $3$  Å (Figure 2).<sup>31</sup>

Do the medium fluctuations described above wash out the influence of the protein fold on ET kinetics as  $R_{DA}$  increases within the database of protein and water-mediated tunneling examined here? If this were the case, in the  $\sigma^2$  dominated regime, all proteins could be described as creating an effective structureless tunneling barrier. To address this central question, which has deep links to the role of evolution in determining coupling pathway structure,<sup>9,33,34</sup> we computed the metric  $S = (\text{var}[\langle H_{DA}^2(R_{DA}) \rangle])^{1/2} / \text{avg}[\langle H_{DA}^2(R_{DA}) \rangle]$ <sup>31</sup> for multi-



**FIGURE 2.** (top) Dependence of  $\ln(C)$  on the DA distance  $R_{DA}$  where  $C = \langle H_{DA} \rangle^2 / \langle H_{DA}^2 \rangle = [1 + (\sigma^2 / \langle H_{DA}^2 \rangle)]^{-1}$ . The dot-dash line denotes the value of  $C$  where  $\langle H_{DA}^2 \rangle = \sigma^2$ . (bottom)  $S = (\text{var}[\langle H_{DA}^2(R_{DA}) \rangle])^{1/2} / \text{avg}[\langle H_{DA}^2(R_{DA}) \rangle]$  (“scatter”) vs distance.  $S$  of the scale of unity indicates that the specific protein structure largely determines the observable rate.<sup>31</sup>

ple ET protein sets characterized by the same transfer distance  $R_{DA}$  (using the Gray–Winkler Ru-proteins<sup>8</sup> as a test set). The variance (var) and the average are taken over all protein ET species at the same transfer distance. We found that  $S$  was of



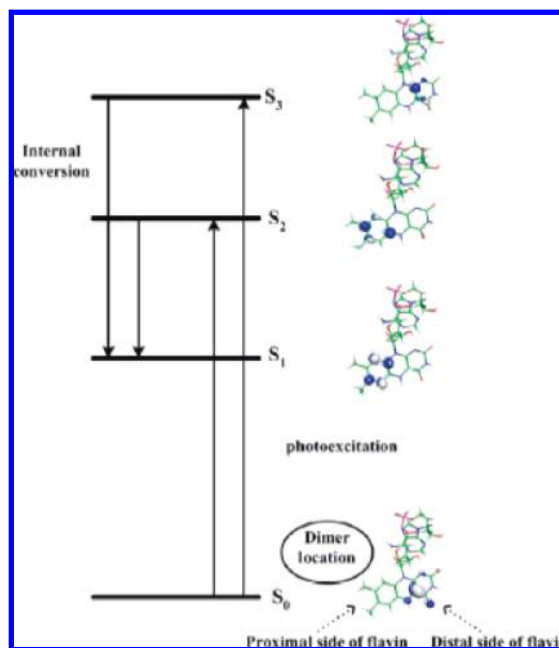
**FIGURE 3.** Mean-squared donor–acceptor interaction as a function of distance between heme edges (larger than the van der Waals gap between protein surfaces) for self-exchange in cytochrome  $b_5$  with protein and water (P,W) produces a coupling plateau compared with the case where only the protein (P) is included.<sup>32</sup>

order unity for all transfer distances (Figure 2, bottom), indicating that the influence of structural differences between proteins on mean-squared ET couplings is large. *Even at large  $R_{DA}$ , where  $\sigma^2$  dominates, the diversity of  $\langle H_{DA}^2 \rangle$  values at any given distance is not eliminated by structural fluctuations.*<sup>31</sup> Thus, the protein structure defines the tunneling rates (for the data set of Ru-proteins examined here) even when fluctuations dominate the ensemble of coupling values. Recent theoretical studies of Ru-proteins<sup>11</sup> and of cytochrome  $bo_3$ ,<sup>21</sup> indicate that the structure of the underlying coupling medium indeed controls the kinetics. Dynamic docking studies of protein–protein ET provide further evidence of the structure sensitivity of ET reactions; these studies find that a very small fraction of the ensemble of docked protein structures is ET active.<sup>35</sup>

### Water-Mediated Electron Tunneling

Water-mediated ET,<sup>36</sup> as noted above, is fluctuation dominated at distances beyond 2–3 Å.<sup>31</sup> Our studies of electron self-exchange in cytochrome  $b_5$  indicate a “coupling plateau” in the distance dependence of water-mediated tunneling.<sup>32</sup> In this plateau regime, structured water bridges between proteins provide a population of strong tunneling routes that include a few water molecules between cofactors (Figure 3). In this regime, structured water bridges establish coupling pathways across otherwise unfavorable van der Waals gaps, enhancing the DA coupling and establishing the plateau.

The model calculations described above provide a framework for understanding an otherwise puzzling array of biological ET studies, including tunneling across water interfaces

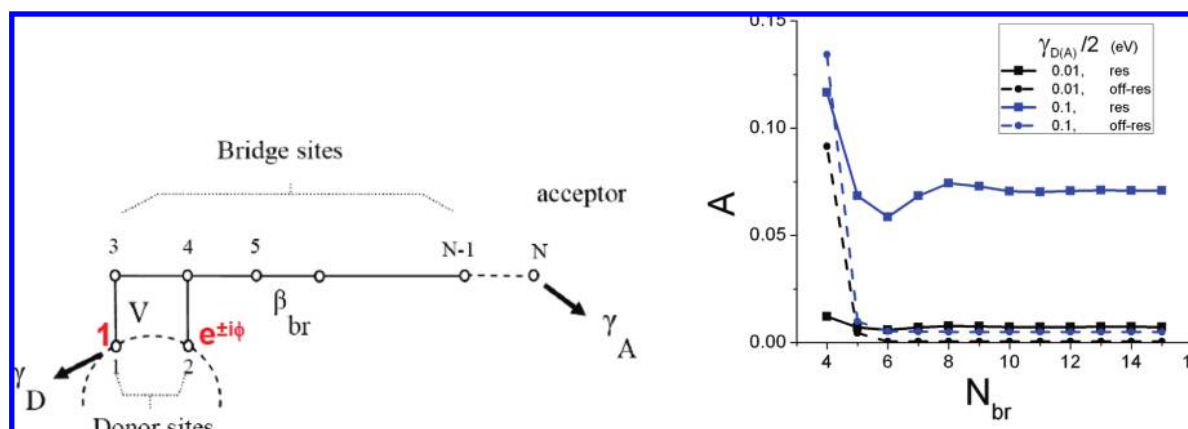


**FIGURE 4.** Electronic energy level diagram for photolyase. The  $S_1$  state is predicted to be polarized strongly in the direction of the thymine dimer.<sup>46</sup>

in the hydroxylating domain of peptidylglycine  $\alpha$ -amidating monooxygenase.<sup>37</sup> Water-mediated tunneling effects have been suggested recently in the Mo-heme ET kinetics of mutant sulfite dehydrogenase<sup>38</sup> and in conducting polymers,<sup>39</sup> as well as in small C-clamp style molecules that were designed to coordinate solvent molecules, including water.<sup>40</sup> Questions of water-mediated tunneling mechanism link to foundational experiments in ET theory (e.g., Fe(II)/Fe(III) self-exchange) that continue to provide test beds for new descriptions of tunneling interactions.<sup>41–43</sup>

### Donor State Polarization and ET Kinetics

In biological ET, a growing body of data indicates that the polarization of the donor state may tune the rate of photoexcited ET. In addition to inducing donor–bridge mixing by the excitation pulse (in the small tunneling energy gap regime<sup>44</sup>), excited-state polarization itself may influence ET kinetics. For example, the excited state of the primary donor in the bacterial photosynthetic reaction center is believed to be strongly polarized in the direction of charge separation.<sup>45</sup> Other ET reactions in photobiology appear to be accelerated by excited states that are polarized toward the acceptor. Our recent theoretical studies<sup>46</sup> of the DNA repair protein photolyase<sup>47</sup> have revealed that photoexcited ET between a reduced flavin adenine dinucleotide ( $FADH^-$ ) and the protein-bound thymine dimer lesion is strongly accelerated by the  $FADH^-$  excited-state polarization in the direction of the dimer lesion.



**FIGURE 5.** A model (left) for ET from angular momenta states D or D\* and plot (right) of the yield asymmetry for D (blue) vs D\* (black) as a function of the number of bridging sites.<sup>52</sup>

We found that the  $\pi-\pi^*$  flavin excited state charge density is polarized strongly toward the acceptor (the “proximal” direction) and that tunneling is dominantly mediated by a bridging methyl group (Figure 4).<sup>46,48</sup> Because of the excited state’s polarization, the nearby adenine, a natural candidate for superexchange mediation due to its unsaturation,<sup>49</sup> appears to play a largely structural role (i.e., a docking site for the thymine dimer), rather than to provide tunneling-mediation pathways.<sup>46,48</sup> The excited-state D polarization helps to explain the fast (subnanosecond) ET from the photolyase excited state to the dimer.

Studies of Skourtis and Nitzan further highlight the potential role of excited-state preparation on ET kinetics.<sup>44</sup> In DNA ET, where the energy gap between D/A and bridge states is small, the initially prepared excited state may contain a considerable admixture of bridge amplitude. Similarly, in molecular conductance experiments, the width of the Fermi–Dirac distribution may populate bridge states, even though the center of the distribution lies in the bridge electronic energy gap. ET in nucleic acids is explored further below.

### Pathway Phase, Chirality, and Electron Transfer

Donor–acceptor interactions are generally computed in the energy domain as the splitting of two quasi-degenerate states.<sup>24</sup> Two interfering coupling pathways with pure real amplitudes of magnitude  $|A|$  and  $|B|$  produce a squared DA interaction proportional to  $(|A| \pm |B|)^2$ , where the + sign enters if the paths interfere constructively and – sign enters if the paths interfere destructively. If the first pathway instead has the complex amplitude  $A e^{i\phi}$ , where  $A$  and  $\phi$  are pure real numbers, the net interaction is proportional to  $A^2 + B^2 + 2AB \cos(\phi)$ . The phase,  $\phi$ , determines the D–bridge interaction, but a simple sign reversal of  $\phi$  has no effect on the magnitude of

the squared DA coupling. In this energy (splitting) framework, the net interaction expected for D and for its complex conjugate D\* is thus identical. As we will see, this energy domain analysis does not indicate that the ET dynamics of D and D\* may in fact be very different in ways that may determine ET rates and yields. A more complete analysis of the electronic dynamics will be required.

Indeed, Waldeck, Naaman, and co-workers showed that experimental ET yields are different for DBA systems with different bridge (B) handedness when the donor state is prepared with circularly polarized light (CPL).<sup>50,51</sup> This result is not explained within the two-state energy-domain analysis above: the energy eigenstate spectra for mirror image systems are essentially identical, and reversing the phase relationship of the D–B interactions (equivalent to reversing the relative phase of the two pathways described in the paragraph above) does not change the magnitude of the coupling. Something is missing from the energy splitting perspective on ET.

The time-dependent transition amplitude from an initial complex (momentum carrying) state D or D\* to a real final state A in a system with Hamiltonian  $H$  is

$$\begin{aligned} \langle A | e^{-iHt/\hbar} | D \rangle &= \sum_m \langle A | \Psi_m \rangle \langle \Psi_m | D \rangle e^{-iE_m t/\hbar} \\ \langle A | e^{-iHt/\hbar} | D^* \rangle &= \sum_m \langle A | \Psi_m \rangle \langle \Psi_m | D^* \rangle e^{-iE_m t/\hbar} \end{aligned} \quad (2)$$

where  $|\Psi_m\rangle(E_m)$  are the eigenstates (energy eigenvalues) of the Hamiltonian. Writing  $\langle A | \Psi_m \rangle \langle \Psi_m | D \rangle = |R_m| \exp[i\Theta_m]$  and  $\langle A | \Psi_m \rangle \langle \Psi_m | D^* \rangle = |R_m| \exp[-i\Theta_m]$ , one finds that the probability of being on the acceptor as a function of time is

$$P_{D,D^* \rightarrow A}(t) = \sum_m |R_m|^2 + 2 \sum_{m>n} \sum_n |R_m| |R_n| \cos[(E_m - E_n)t/\hbar \mp (\Theta_m - \Theta_n)] \quad (3)$$

Importantly, switching the donor state  $D^*$  to (accomplished by reversing the polarization of the CPL) introduces phase shifts in the time evolution. Reversing the helicity of a bridge that interacts with a ring-like donor state has an equivalent effect. Even in achiral systems, bridges that interact with the complex phase donor at more than one site will display different time evolutions for the D to A vs the  $D^*$  to A transition probabilities.<sup>52,53</sup>

Introducing D and A state lifetimes in eq 3 ( $\hbar/\gamma_D, \hbar/\gamma_A$ ) produces phase-dependent ET yields,  $Y(D) = \gamma_A \int_0^\infty dt P_{D \rightarrow A}(t)$ , and a yield asymmetry:

$$A \equiv \frac{Y(D^*) - Y(D)}{Y(D^*) + Y(D)} \quad (4)$$

This predicted asymmetry was validated in tight-binding calculations for the model of Figure 5,<sup>52</sup> which produces yield asymmetries consistent with the experimental results.<sup>50,51</sup> Figure 5 shows that the yield asymmetry grows with  $\gamma_D$  (zero when  $\gamma_D = 0$ ) because donor relaxation “locks in” the difference in the propagation from the prepared states of opposite phase.

It is remarkable that changes in the initial state orbital angular momentum or of the bridge handedness introduce detectable differences in ET yields. Introducing donor state angular momentum necessitates introducing complex amplitudes, and bridges that interact with the complex donor state at more than one site produce pathway interferences that reflect the phase relationship among donor sites. It will be fascinating to expand this framework for describing “momentum transfer” to the regime of molecular junctions, as well as to unimolecular ET kinetics.

## A Molecular Double-Slit Experiment

In the multipathway ET systems discussed above, the electron propagates coherently from D to A. That is,  $H_{DA}$  is computed by adding the amplitudes arising from all bridging pathways. The coherent donor–acceptor interactions mediated by a bridge can be written as a sum over products of amplitudes propagating among chains of orbitals with interaction  $V_{nm}$ . This sum is known as a Dyson expansion and carries the McConnell model<sup>15,24,54</sup> for DA interactions to infinite order. The lowest-order contribution to the coupling along pathway  $n$  from D to A is

$$h_{DA}^{(n)}(\text{Path}[D \rightarrow i \rightarrow j \rightarrow k \rightarrow \dots z \rightarrow A]) = \frac{V_{Di}V_{ij}V_{jk} \dots V_{zA}}{(E_i - E_j)(E_j - E_k)(E_k - E_l) \dots (E_z - E_A)} \quad (5)$$

and the total coupling (summed coherently over all pathways) is

$$H_{DA}^{\text{elastic}} = \sum_n h_{DA}^{(n)} \quad (6)$$

where  $E_i$  is the tunneling energy of the electron,  $E_i$  is the energy of orbital  $i$  along the pathway, and there is no energy exchange between the electron and bridge vibrations.

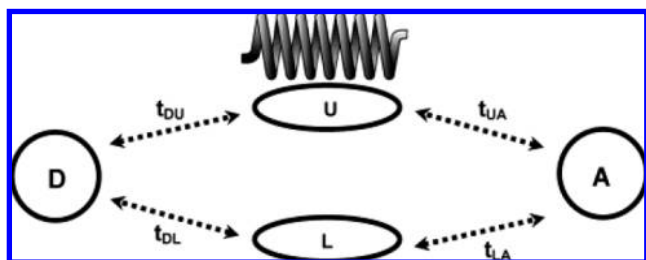
Imagine that when the electron occupies orbital  $j'$  in one of the tunneling pathways contributing to the sum, it interacts and exchanges energy with a localized vibrational mode (e.g., a carbonyl vibration) enabling inelastic tunneling. If the electron tunneling event changes the quantum vibrational state of the mode localized on orbital  $j'$ , that vibrational excitation is evidence that the electron visited orbital  $j'$  while tunneling. Thus, only pathways in the summation of eq 6 that include orbital  $j'$  (where energy is exchanged between the electron and nuclear vibrations) enter the DA coupling (eq 7), in contrast to the elastic case where all pathways enter the summation:<sup>55–57</sup>

$$H_{DA}^{\text{inelastic}} = \sum_n h_{DA}^{(n,\text{inel})}(j') \quad (7)$$

Since pathways may interfere constructively or destructively, inelastic events can result in either increasing or decreasing  $H_{DA}$ . The inelastic effect described here arises for the same reason that detecting the electron's path in a double-slit experiment destroys interference between alternative paths.<sup>10</sup> In ET molecules, this inelastic effect may be particularly dramatic if the elastic coupling pathways interfere destructively (as in orbital symmetry-forbidden ET). As such, turning on an inelastic transport mechanism may switch on an otherwise forbidden kinetic process. This effect is different from other effects of inelastic interactions on the tunneling matrix element and on the nuclear Franck–Condon factor, which have been studied.<sup>18,58</sup>

The inelastic tunneling pathway effect described here is only predicted when a fully quantum treatment is made of bridge-mediated inelastic tunneling (i.e., when coupling pathways are defined as sequences of vibronic states, that are products of electronic orbitals and bridge vibrational states).<sup>55–57</sup>

For the model in Figure 6, to lowest order in the electronic coupling interactions  $t_{ij}$ , the elastic coupling is  $H_{DA}^{\text{elastic}} \approx (t_{DU}t_{UA}/\Delta E) + (t_{DL}t_{LA}/\Delta E)$  ( $\Delta E$  is the D/A to bridge orbital electronic energy gap), and the inelastic coupling is  $H_{DA}^{\text{inelastic}} \approx (\gamma t_{DU}t_{UA}/\Delta E^2)$  (where  $\gamma$  is the electron-vibrational coupling on the bridge). The inelastic path coupling thus has only one bridge pathway contribution, while the elastic coupling has two parallel path contributions. The inelastic path is expected to be weaker by about 1 order of magnitude compared with the



**FIGURE 6.** A model DBA system with vibronic coupling to the upper bridge orbital. D, U, L, and A denote donor, bridge, and acceptor electronic states, respectively. The spring represents a bridge-localized vibration that is perturbed when the electron visits orbital U.<sup>55–57</sup>

elastic path in many systems.<sup>55</sup> As such, inelastic effects will be most pronounced in systems with symmetry-forbidden elastic ET kinetics that will be allowed in the inelastic regime.

A detailed quantum analysis of intramolecular tunneling in more realistic model molecular system confirms the argument that inelastic tunneling via localized vibrational modes will allow the selection of coupling pathways.<sup>57</sup> Indeed, we have proposed a scheme for driving local normal modes with IR radiation, thus promoting inelastic tunneling,<sup>57</sup> and experimental tests of this conjecture are in progress. Pathway manipulation via inelastic ET provides an example where *bridge vibrational dynamics* enables control of the bridge-mediated donor–acceptor electronic communication and, hence, of ET kinetics. The goal of designing, synthesizing, and experimentally probing unimolecular inelastic tunneling systems (with

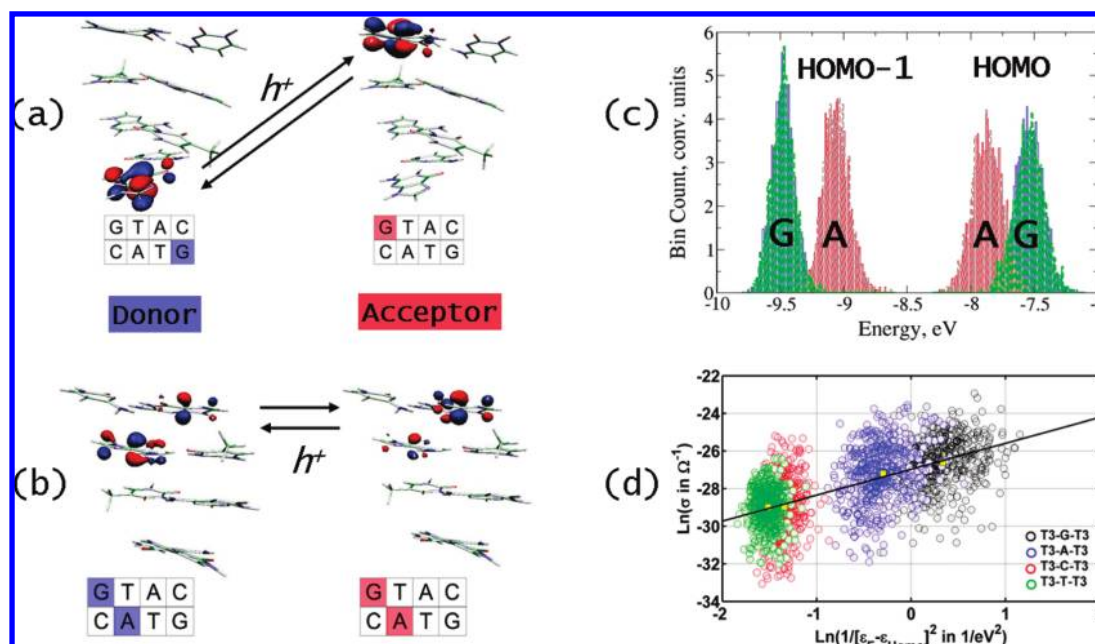
spatially localized vibrational modes) represents a grand challenge.<sup>3,27,59–61</sup>

## ET in Nucleic Acids and the Effects of Thermal Fluctuations

Conformational fluctuations play a central role in nucleic acid (NA) ET. The accessibility of many viable transport mechanisms, including superexchange, incoherent multistep hopping, and polaronic transport, rely upon medium fluctuations (see ref 62 and references therein). The reported distance dependences of ET in NAs, from nearly distance-independent to exponentially decaying, are linked both to the average energetics of the donor, bridge, and acceptor states and to fluctuations in their energies and couplings (Figure 7a–c).<sup>62</sup>

In the deep-tunneling regime, (accessed when the donor/acceptor energies are far from the base energies), the thermally averaged ET rates decay approximately exponentially with distance, and fluctuations of base energetics and interbase couplings cannot provide access to hopping or other transport mechanisms. Molecular conductance calculations performed on single stranded NA fragments show that, despite a significant spread in conductance values arising from fluctuations in base energies and couplings, a McConnell superexchange model based upon average nucleobase energies qualitatively describes the ET (Figure 7d).<sup>63</sup>

In contrast to the case of the deep tunneling regime (Figure 7d), NA ET can also occur in systems with relatively small



**FIGURE 7.** (a) G-localized hole states of the terminal Gs in the CATG DNA segment.<sup>64</sup> (b) Hole states shared between G and A. (c) Histogram of fluctuating HOMO and HOMO – 1 energies. Note the significant overlap of the G and A orbital energies. (d) The conductance ( $\sigma$ ) computed for individual snapshots of the single-strand PNA segment TTTXTTT ( $X = A, C, T, G$ ) shows scatter over orders of magnitude because of fluctuation in both base energies and interbase couplings.<sup>63</sup>

redox potential gaps between the donor/acceptor and bridge states, leading to a mixed mechanism (see Figure 7a–c). The influence of fluctuations on ET kinetics and mechanism becomes very different as the donor/acceptor–bridge barrier height shrinks, as in the case of G–G hole transport across NAs (Figure 7).<sup>64</sup> For a G base bound hole, the energy gap to the nearest bridge hole states (A) is on the scale of tenths of electronvolts for the bases in their average geometries. As a consequence, most ET models assume a tunneling mechanism for short distance transfer and a hopping mechanism over larger distances (typically, above three to four base pairs). Indeed, tunneling–hopping transitions are observed in conjugated polymers<sup>65</sup> as a function of length, and the crossover presumably arises when the propensity for carrier injection into bridge states exceeds the tunneling probability. Our recent simulations, however, indicate that the two mechanisms may not be so clearly separated in NA ET.<sup>64</sup> ET theory predicts electronic energy fluctuations on the scale of  $\delta E \approx (\lambda k_B T)^{1/2}$  where  $\lambda$  is the reorganization energy. For NAs,  $\lambda \approx 1$  eV so  $\delta E \approx 0.1$ – $0.2$  eV at room temperature, which is on the same scale as the difference in the oxidation potentials for G and A bases.<sup>64</sup> Indeed, a 0.1–0.2 eV thermal spread in frontier orbital energies was computed in our recent analysis of the electronic structure fluctuations for short NA segments (Figure 7c).<sup>64</sup> The principal fluctuations leading to the energy spread appear to arise from covalent bond length changes of the individual bases, rather than from fluctuations of NA helical parameters.

Thermal fluctuations thus bring the G and A bases into frequent degeneracy (see Figure 7), despite the fact that their redox potentials differ by ten times thermal energies or more. The resulting hole delocalization between the G and A (Figure 7a,b) indicates a possibility of hopping transport even at short G-to-G distances. In our analysis of the CATG DNA/PNA sequence, we found about 10% of the structural snapshots displayed significant hole delocalization away from the Gs.<sup>64</sup> These studies suggest that fluctuations may limit the viability of pure superexchange or pure hopping models for hole transfer in NAs, even at short distances. Previous theoretical studies (see refs 66 and 67 and references therein) have shown that, in sufficiently low gap ET systems, the tunneling matrix element and the two-state golden rule description of ET are not appropriate for describing the observed kinetics because of degeneracy, or near degeneracy, with bridge states. It will be interesting to reexamine kinetic data associated with DNA and PNA hole transport in light of such mixed mechanisms.

## Conclusions and Prospects

In proteins, the motif and ET distance determine whether fluctuations or average structure control the coupling mechanism. Water structure and fluctuations add additional surprises; organized water between proteins in near contact appears to provide enhanced tunneling propagation despite the large bridge orbital energy gap for water. Excited-state preparation also influences ET kinetics and dynamics. Donor states with different angular momenta propagate differently, as a consequence of the complex valued donor state amplitudes, giving rise to new ET effects that are not predicted by analyzing energy splittings. Further, polarized excited donor states were shown to select specific coupling pathways in proteins and to have large effects on ET rates, especially for large donors. This principle is particularly important in the protein photolyase, where polarization of the donor appears to accelerate DNA repair, likely enhancing the protein's function.

DNA and PNA provide examples where fluctuations of bridge energetics influence ET coupling interactions as well as the transport mechanism. Since energy fluctuations are on the same scale as the differences among the redox potentials of the bases, fluctuations will likely mix superexchange, hopping, and polaronic transport regimes.

Finally, a perspective that includes inelastic coupling pathways may provide schemes to control ET. Indeed, leaving behind vibrational “breadcrumbs” that label tunneling pathways should change the donor–acceptor interaction and the ET kinetics. It seems likely that schemes being developed to control coherence and entanglement in meso- and nanoscale quantum structures will find intriguing mappings in the world of molecular ET.

*The authors thank the National Institutes of Health, the National Science Foundation, the Department of Energy, and the University of Cyprus for support of our research programs. We are grateful to our students and postdocs, Horacio Carias, Liz Hatcher, Tsutomu Kawatsu, Igor Kurnikov, Jianping Lin, and Tatiana Prytkova, and to our collaborators, Catalina Achim, Harry Gray, Brian Hoffman, Fred Lewis, Marcela Madrid, Ron Naaman, Abe Nitzan, José Onuchic, Igor Rubtsov, Jon Sessler, Dave Waldeck, and Jay Winkler, for their enthusiastic interactions on these projects.*

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