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Mutual Orientation Effects on Electron Transfer between Porphyrins

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Mutual orientation effects on the rate of nonadiabatic electron transfer between several diporphyrin pairs of experimental interest are examined. The electronic matrix element for electron transfer is calculated within a one-electron spheroidal model for a variety of states and orientations which are relevant to both biological and synthetic electron-transfer systems. Both the mutual orientation of the pairs and the nodal structure of the donor and acceptor orbitals can have large effects on calculated rates.

I. Introduction

As increasingly detailed information is obtained on the rates and mechanisms of electron-transfer reactions, it has become evident that a more complete understanding is desirable of how the separation distance and mutual orientation of a reactant pair affect the rate. This is especially true for biological and biomimetic electron-transfer pairs. For example, cytochrome c is a widely studied biological electron-transfer agent. The heme is oriented with respect to the protein and has one edge exposed to the solution environment. Its reactions have been discussed in this context. Many other biological electron transfers occur between molecules held rigidly at fixed distances and orientations. The electron-transfer rate has been measured for a [ZnFe] hybrid hemoglobin, for which both the orientation and separation distance of the two porphyrins are known. Relative orientations of biological electron-transfer pairs are under study: It has been reported that the molecular planes of the heme c and d groups in cytochrome c-d of Pseudomonas aeruginosa are perpendicular to one another in both the reduced and oxidized forms of the protein. Also, the relative orientations of the initial charge-transfer agents of photosynthetic reaction centers have been determined. Synthetic systems have also exhibited possible orientation-dependent electron transfers. Photoexcited electron transfer has been observed in the cofacial diporphyrins of Chang, and it has been found to proceed rapidly in the forward direction and considerably slower in the reverse. Systems that are similar, but where the transition moments of the two porphyrins subunits are oriented perpendicularly, rather than parallel, have been examined by Overfield et al. They show much slower forward transfer than those of ref 7-10. Separation-distance effects have been examined in rigidly linked porphyrin–quinone systems, so chosen because biological electron transfers frequently involve aromatic donors and acceptors such as porphyrins, porphyrin derivatives, and quinones. In addition, charge-transfer and crystallographic studies have been made of another set of donor–acceptor pairs linked as metacyclophanes and paracyclophanes. Also, it has been proposed that electron-transfer fluorescence quenching of chlorophyll a by quinone multilayer arrays requires a favorable orientation of reactants and products.

Orientation effects have been studied theoretically by using a variety of models to examine qualitative effects. Orientation effects have also been examined in studies of the ordering of spin states in compounds containing two metal centers, a problem analogous to intramolecular electron transfer. We have recently described a simple model for orientation effects based on the delocalized nature of σ-electrons in aromatic systems. It is a one-electron model in which the transferable electron is assumed to be bound at an oblate-spherical potential well of specified depth. Electron transfer was modeled between two such nonpenetrating sites, and the effects of orientation on the thermal matrix element (the electronic matrix element appearing in theories of nonadiabatic electron transfer) were examined for various separation distances.

In the present paper the model of ref 27 is applied to a number of systems of experimental interest, both biological and synthetic. The aim of the model is to illustrate possible effects of orbital and...
potential shape on the rate of electron transfer at fixed distances and orientations. Such calculations, to the extents that they are applicable, can reveal ways in which the architecture of individual electron-transfer pairs may help control electron-transfer rates. In particular, orientation effects are examined for the forward reaction \( A^+ + B \rightarrow A^+ + B \) for several systems, \( A^+ \) denoting a photoexcited molecule, and for the highly exothermic reverse reaction \( A^+ + B^- \rightarrow A + B \). The systems for which calculations are reported are (a) synthetic face-to-face porphyrins, including an open-jawed configuration, (b) porphyrin-like systems in an edge-to-edge configuration, (c) porphyrin systems comparing edge-to-edge and face-to-face as well as intermediate configurations, and (d) a photosynthetic system involving a dimeric photoexcited chlorophyll molecule.

The actual experimental results for which orientation effects have been explicitly studied are at present relatively few. In particular, there are the results of Chang et al.\textsuperscript{7-10} and Overfield et al.\textsuperscript{11,12} which are compared with the present results below. The present article is organized as follows: The model is summarized and the methods of choosing states and energies for given systems are discussed in section II. Calculated results for several physical systems are given in section III, and they are discussed in section IV.

\section*{II. Theoretical Model}

In current theories of nonadiabatic electron transfer,\textsuperscript{28-30} the rate constant for electron transfer between two reactants, A and B, at a given fixed separation distance and orientation is given by

\[ k_{B-A} = \frac{2\pi}{\hbar^2} |T_{BA}|^2 FC \]

(1)

In eq 1, FC is a sum of thermally weighted Franck–Condor factors for the nuclear vibrational, librational, and rotational coordinates of the two reactants and the surrounding medium.\textsuperscript{28-34} The distance and the orientation dependences of the rate constant occur mainly in the factor \( T_{BA} \) given by\textsuperscript{27,29,35}

\[ T_{BA} = (H_{BA} - S_{AB} H_{AA})/(1 - |S_{AB}|^2) \]

(2)

where

\[ H_{BA} = \langle \Psi_B | H_{A} | \Psi_A \rangle, \quad H_{AA} = \langle \Psi_A | H_{A} | \Psi_A \rangle \]

(3)

\[ S_{AB} = \langle \Psi_A | \Psi_B \rangle \]

(4)

With this Hamiltonian an energy \( E \) can be computed, given by \( \langle \Psi_A | H_{A} | \Psi_A \rangle \). For the given value of \( H_{A} \), this \( E \) becomes a vertical one-electron ionization potential from the orbital \( \Psi_A \).

Exact (three-dimensional) eigenfunctions were calculated for this Hamiltonian. Oblate-spheroidal coordinates\textsuperscript{36} (\( \xi, \eta, \varphi \)) (Figure 2) were used to solve the Schroedinger equation. The coordinate \( \varphi \) is the angle of rotation around the \( z \) axis. The wave function \( \Psi(\xi, \eta, \varphi) \) can be written as \( \Psi(\xi, \eta) \Psi(\varphi) \). The Schroedinger equation can then be separated with respect to \( \varphi \), and one obtains \( c_1 \cos m\varphi + c_2 \sin m\varphi \) for \( \Psi_{m}(\varphi) \). The separation constant \( m \) depends on the given quantum state of interest. The function \( \Psi(\xi, \eta) \) is even or odd with respect to reflection in the \( xy \) plane. There are three types of possible nodal surfaces for \( \Psi \), roughly corresponding to constant-coordinate surfaces for the three coordinates \( \xi, \eta, \) and \( \varphi \). An \( \xi \)-type nodal surface is radial-like. (More precisely, at large distances \( d/2 \) is approximately equal to the distance from the center of the spheroid.) Wave functions with one \( \eta \)-type node have, by symmetry, the \( xy \) plane of the potential as a nodal surface; i.e., they have \( \eta \)-symmetry with respect to the \( xy \) plane. The \( \varphi \)-like nature of such states is shown in Figures 3a and 4a. Higher numbers of \( \eta \)-type nodes are symmetrically placed about the \( xy \) plane. The \( \varphi \)-type nodal surfaces are planes through the origin, perpendicular to the molecular \( xy \) plane. Contour plots of states having \( m = 4 \) and \( m = 5 \) are given in Figures 3 and 4. In particular, Figures 3b and 4b are useful in visualizing the \( \varphi \)-type nodal surfaces.

When two oblate-spheroidal potential wells are chosen at a given separation distance and orientation, each with a specific single-site wave function, the various integrals in eq 3 and 4 can be evaluated

\[ H_{A} = -\frac{1}{2} \nabla^2 + V_{A} \]
The present results given below is in terms of the three-dimensional expression. Accordingly, only values of the dimensional integral, and for that reason, the discussion of the results given below is in terms of the three-dimensional expression. The states chosen for a given calculation of $H_{BA}$ are dependent upon the molecular system being modeled. Since the present work by Gouterman et al., united the ring model description with the molecular orbital approach in what has become known as the “four-orbital model.”

Figure 3. Contours of $\Psi$ for a $(4,\pi)$ state for $V_0 = 19.2227$ eV, $E = -0.4000$ eV, $a = 5$ Å, and $b = 2$ Å. The heavy line is the well boundary. The contours are labeled with values of $\log |\Psi|$. Dashed lines indicate $\Psi < 0$, and solid-line contours are for $\Psi > 0$. In (a) the contours shown are in the $xz$ plane. In (b) the contours shown are in a plane parallel to the $xy$ plane which intersects the $z$ axis at $z = 1$ Å.

Figure 4. Contours of $\Psi$ for a $(5,\pi)$ state for $V_0 = 23.4040$ eV, $E = -0.4000$ eV, $a = 5$ Å, and $b = 2$ Å. The labeling conventions of Figure 3 were followed.

The article is concerned with electron transfers involving porphyrins and related compounds, the method of selecting the states appropriate to these systems is discussed. In all cases it is assumed that the transferable electron is delocalized over the porphyrin ring and does not have significant density on any central metal atom. A similar rationale could be applied to other cyclic aromatics.

In one early theoretical attempt to understand porphyrin spectra the $\pi$-electrons were treated as being confined to a one-dimensional ring.

Due to the definition of $V_B$, the expressions in eq 3 then reduce to

$$H_{BA} = -V_0 <\Psi_B|\Psi_A> B, H_{AA} = -V_0 <\Psi_A|\Psi_A> B$$

where the subscript B on the integrals indicates that integration is only over well B. For the states used in the present article it was found in all cases that $T_{BA}$ was equal to $H_{BA} to within 3\%$ when $H_{BA}$ was nonzero and that the accuracy increased with separation distance. Furthermore, for a pair of $(4,\pi)$ states having the energy used in ref 27, it was found that the zeros of $T_{BA}$ and $H_{BA}$, for the orientations of Figure 5d, were within two degrees of one another when the wells were in contact. Accordingly, only values of $H_{BA}$ are presented in this paper.

In Appendix A a method is shown for converting $H_{BA}$ to a two-dimensional integral which reduces the computation time for $H_{BA}$ significantly. The numerical results are the same as those from the direct three-dimensional integration over well B and were used here. Conceptually, it is easier to envision the three-dimensional integral, and for that reason, the discussion of the results given below is in terms of the three-dimensional expression.

The states chosen for a given calculation of $H_{BA}$ are dependent upon the molecular system being modeled. Since the present article is concerned with electron transfers involving porphyrins and related compounds, the method of selecting the states appropriate to these systems is discussed. In all cases it is assumed that the transferable electron is delocalized over the porphyrin ring and does not have significant density on any central metal atom. A similar rationale could be applied to other cyclic aromatics.

In one early theoretical attempt to understand porphyrin spectra the $\pi$-electrons were treated as being confined to a one-dimensional ring.37 Within this treatment, the eigenfunctions are of the form $\exp(\pm im\phi)$, the highest occupied pair of orbitals having $m = 4$ and the lowest unoccupied pair having $m = 5$.37 Later theoretical work by Gouterman38,39 united the ring model description with a molecular orbital approach in what has become known as the “four-orbital model.”

The wave functions obtained in the molecular orbital approach are real, the HOMO and HOMO resembling the cos $4\phi$ and $4\phi$, and the LUMO and LUMO resembling the cos $5\phi$ and $5\phi$. Later ab initio calculations31-34 have further supported


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In this article, states with one \( \pi \)-type node (which are \( \sigma \)-states) and with \( m = 4 \) or 5 (designated \( \psi_4 \) and \( \psi_5 \), respectively) were chosen to qualitatively reproduce the four-orbital model states. Figures 3 and 4 show contour plots of \( \psi_4 \) and \( \psi_5 \) states. The nodal patterns are qualitatively similar to those of the HOMO and LUMO orbitals in ab initio calculations for porphyrin-like molecules. Metallo- and free-base porphyrins, chlorophylls, and bacteriochlorophylls were all treated as having the same HOMO's and LUMO's.

In modeling the first excited singlet state of a given molecule, one only considers the excited, transferable electron. The four-orbital model predicts that the first excited singlet state is composed of linear combinations of pairs of the possible primitive excitations formed from the four orbitals of interest. The extent of mixing of the possible single excitations will determine the fraction of mixing of \( \cos \psi \) and \( \sin \psi \) in the donor wave function (cf. ref 38, 39, and Appendix B). This mixing will depend on the symmetry, substituents, and the environment of the molecule.

Using the four-orbital model and semiempirical or ab initio molecular orbital calculations, we could estimate the extent of mixing between the pair of single excitations which make up the first excited singlet state, thus yielding an approximate description of a given excited state within the present model (see Appendix B). However, the results below are presented for \( \Phi_m(\psi) = \cos \psi \) or \( \sin \psi \) to ensure that the orientation effects seen are not peculiar to a specific choice of \( \Phi_m(\psi) \). Any \( \Phi_m(\psi) \) can be generated as a linear combination of the above.

To represent the HOMO of a (metal-)porphyrin anion \( \Phi_m = \cos \psi \) or \( \sin \psi \) was again chosen, on the assumption that the additional electron is placed in the LUMO of the neutral molecule. ESR data on bacteriochlorophylls (BChl) and bacteriopheophytins (BPh) indicate that the unpaired spin density is delocalized over the entire ring. The spin densities obtained from ab initio and semiempirical calculations are in reasonable agreement with experimental results and can be approximately described as having the extra electron in the LUMO of the neutral molecule.

Finally, in modeling the empty orbital in (metallo)porphyrin cations, we selected \( \Phi_m(\psi) = \cos \psi \) or \( \sin \psi \) states. Evidence from comparisons of MO calculations with ESR measurements on porphyrins suggests that these cations can be described qualitatively by such localized, single-electron hole descriptions. In the case of certain metalloporphyrins only a few percent of the electron density is delocalized on the porphyrin ring. The present results are not applicable, therefore, to transfers through heme faces. It is possible that the calculations are still applicable to transfer through heme edges, but with appropriately reduced values of the matrix elements \( T_{BA} \).

In summary, in the results of the following section \( H_{BA} \) is examined as a function of orientation for transfer of an electron initially localized in a \( \psi \) state and for transfer of an electron initially localized in a \( \psi \) orbital. The \( \psi \) orbital can be viewed as a forward transfer between a reduced electron and a neutral acceptor. The \( \psi \) transfer can be viewed as a back transfer from a reduced acceptor to an oxidized donor to yield the ground-state species.

The sizes of the potential wells were chosen as follows. The semimajor axis \( a \) was chosen as 5 Å, an approximate in-plane radius of the porphyrine system. The semiminor axis \( b \) was chosen as 2 Å. This \( b \) yields an average thickness of 2.7 Å for the spheroidal well. This value represents approximately the "thickness" of the electron cloud obtained from ab initio calculations on a substituted porphyrin and allows approach of the "molecular" planes to distances close to those found in synthetic porphyrins.
systems. As in ref 27, the trends in the results presented on orientation dependence are not sensitive to the exact values of $a$ and $b$.

The energy $E$ was chosen to make the decay of $H_{BA}$, with separation distance, lie in the general vicinity of a number of experimental estimates. In ref 27 it was shown that $H_{BA}(R)$, in the present model decayed approximately exponentially with increasing separation of the wells. The calculated decay constant $\beta$ for $\ln |H_{BA}|^2$ depends on the range of distances for $H_{BA}$ being used, since $H_{BA}$ is not a pure exponential. Here, the estimated $\beta's$ were obtained for edge-to-edge separations of between 10 and 20 Å since several experimental studies have produced estimates of $\beta$ for transfers at these distances. (Such distances are appropriate to those estimated in studies of tunneling in glassy matrices.) In general, the calculated $\beta$ decreases with increasing distance between the wells. The calculated $\beta$ also depends on the orientation of the wells. The orientations chosen for estimating $\beta$ were those of Figure 5d. The angles $\Theta = 0^\circ$, $60^\circ$, and $90^\circ$ were chosen as representative; $\Theta = 60^\circ$ approximately corresponds to the maximum in $H_{BA}$ as a function of $\Theta$ for a given edge-to-edge distance for this class of orientations. We have chosen $E = -0.400$ eV which, for $(5,\pi) \rightarrow (5,\pi)$ transfer, yields $\beta's$ of 1.5, 1.4, and 1.3 Å$^{-1}$ for $\Theta = 0^\circ$, $60^\circ$, and $90^\circ$, respectively, over the above range of distances. Since $\Theta = 60^\circ$ corresponds to the maximum in $H_{BA}$ for the class of orientations, its decay should be most important in determining $\beta$ when averaged over $\Theta$. Other orientations may yield different $\beta's$, but they are expected to fall in the range of the above values.

In electron-transfer reactions, the energy region of interest is that near the intersection of the reactants’ and products’ potential energy surfaces, i.e., at the intersection state for the reaction. In this region the reactants’ and products’ electronic energies are equal. The donor and acceptor potential depths were thus adjusted to make the energies of $\Psi_A(E) = (\Psi_A | H_{BA} | \Psi_A)$ and $\Psi_B(E) = (\Psi_B | H_{BA} | \Psi_B)$ equal to $-0.400$ eV, for both forward and reverse transfer. The relative angular dependence is largely unaffected by the orbital energies used.

III. Results

Results are presented in this section for the set of orientations depicted in Figure 5. In Figure 5a the $xy$ planes of the two wells are assumed parallel, and $\delta$ is the distance between the $z$ axes of the two wells along their common $x$ direction. The orientations in Figure 5b involve a rocking motion of the two porphyrins through the angle $\alpha$, the $xy$ planes of the wells being parallel at $\alpha = 0^\circ$. The interplane separation at the assumed hinge point is fixed at the value $\Delta z$. The orientations described in Figure 5c involve an edge-to-edge configuration of the wells. The wells are first translated relative to one another along their common $x$ axis, and then one well is tilted through an angle $\Gamma$ as shown in Figure 5c. The edge-to-edge distance along the common $x$ axis for the orientations of Figure 5c is denoted by the parameter $d$ (not shown). The orientations in Figure 5d involve wells with parallel $xy$ planes, with the origin of the second well moved through the swing angle $\Theta$. The parameter $\sigma$ is the edge-to-edge separation distance. In Figure 5e, a set of orientations used to examine possible electronic effects in bacterial photosynthetic reaction centers is shown.

Values of $H_{BA}$ are plotted in Figures 6 and 7 for the states used were as in Figure 6. For back transfer $(5,\pi) \rightarrow (4,\pi)$, the states used were as in Figure 7. In both cases initial and final states were of the form $\cos mc$. The labeling convention of Figure 6 was used.

![Figure 6](image_url)

**Figure 6.** Matrix element $H_{BA}$ as a function of the displacement parameter $\delta$ defined in Figure 5a at two fixed interplane separations for $(5,\pi) \rightarrow (5,\pi)$ transfer: (a) interplane spacing = 4.2 Å; (b) interplane spacing = 5.4 Å. For the donor and acceptor states $\alpha$, $\delta$, and $E$ were as in Figure 4. The solid line indicates $\Psi_A(E) = \cos mc$ in each well; the dashed line indicates $\Psi_B(E) = \sin mc$ in each well.

![Figure 7](image_url)

**Figure 7.** Matrix element $H_{BA}$ as a function of the displacement parameter $\delta$ defined in Figure 5a at two fixed interplane separations for $(5,\pi) \rightarrow (4,\pi)$ transfer: (a) interplane spacing = 4.2 Å; (b) interplane spacing = 5.4 Å. For the donor and acceptor states $\alpha$, $\delta$, and $E$ were as in Figure 4. For the donor, $V_0$ was as in Figure 4, and for the acceptor, $V_0$ was in Figure 3. The labeling convention of Figure 6 was used.

![Figure 8](image_url)

**Figure 8.** Matrix element $H_{BA}$ as a function of the rocking angle $\alpha$ defined in Figure 5b between the two porphyrin planes. For forward transfer $(5,\pi) \rightarrow (5,\pi)$ the states used were as in Figure 6. For back transfer $(5,\pi) \rightarrow (4,\pi)$, the states used were as in Figure 7. In both cases initial and final states were of the form $\cos mc$. The labeling convention of Figure 6 was used.
IV. Discussion

The results of the previous section are considered here in the order presented there.

(56) On the basis of recent optical evidence [Meech, S. R.; Hoff, A. J.; Wiersma, D. A. Chem. Phys. Lett. 1985, 121, 287–292] it has been proposed that the initial excitation of the BChl dimer is followed immediately (~0.025 ps) by the formation of an intramolecular charge-transfer state of the dimer. Subsequent charge transfer, e.g., to the BChl monomer, could then involve a transfer from the anion in the dimer.
avoided crossing of the electronic surfaces, which can occur in forward and reverse directions, this avoided crossing in the inverted region will not cause any greater decrease in the rate than does the nonadiabaticity (i.e., the small activation energy of the process, Le., the ‘inverted effect”)

The results of Figures 6 and 7 indicate that an orbital orientation effect could also contribute to the difference in rates in this face-to-face configuration.

The existence of a zero of \( H_{BA} \) at \( \delta = 0 \) is purely an orbital shape effect, due to the orthogonality of the \( \Phi_m(\phi) \) functions for \( m = 4 \) and \( m = 5 \) in the face-to-face configuration. When the \( z \) axes of the two wells lie along a common line, the product \( \Phi_s(\phi_A) \Phi_s(\phi_B) \) vanishes by symmetry when integrated over \( \phi_A \). (Note that the integration is over well \( A \) in this case since back transfer occurs from \( B \) to \( A \).) This large difference in forward and reverse matrix elements is very orientation dependent: Comparing Figures 6 and 7, it is seen that the forward and back transfer \( H_{BA} \)'s become of comparable magnitude for \( \delta > 3 \) \( \text{Å} \). The existence of a zero of \( H_{BA} \) at \( \delta = 0 \) applies also, exactly, to \( T_{BA} \), as one can show by a symmetry argument. The same remark applies for the back transfer results at \( \alpha = 0 \) in Figure 8 given below.

Experimental results on such face-to-face porphyrins appear to indicate fast, light-driven forward electron transfer (<6 ps) with slow back transfer to yield ground-state products (~1 ns). It has been suggested that the back transfer is slow due to a large avoided crossing of the electronic surfaces, which can occur in the inverted region, or because of the large driving force and small activation energy of the process, i.e., the “inverted effect”, or both. The results of Figures 6 and 7 indicate that an orbital orientation effect could also contribute to the difference in rates in this face-to-face configuration.

The results given in Figure 8 show a dramatic dependence of \( H_{BA} \) on the jawing angle \( \alpha \) for face-to-face porphyrins. For the \((5,\pi) \rightarrow (5,\pi)\) and \((5,\pi) \rightarrow (4,\pi)\) transfers it is seen that only for values of \( \alpha \) less than 20° are the forward and back transfer \( H_{BA} \)'s significantly different. For larger jawing angles the forward and back transfer \( H_{BA} \)'s are virtually the same.

While there are no experimental results for the jawed configurations in Figure 5b, there are the results of Overfield et al.11,12 for a somewhat different set of jawing configurations. In these studies evidence was obtained that suggests that forward and back transfer occur at essentially the same rate in the jawing diporphyrins studied, each state having a lifetime of ~1 ns. This result is quite interesting because the interplane separation in this compound is comparable to that in the compounds of Chang,10 and for small jawing angles, the molecules are planar. Such slow forward and back transfer may also be due to an orbital orientation effect.

The compound examined by Overfield et al.11,12 is a jawing diporphyrin, similar to that of Figure 5c, except that one of the wells is rotated 180° about a line in the \( xy \) plane of the potential which bisects the angle between the \( +x \) and \( +y \) axes. (Thus, the \( x \) axes of the wells are antiparallel, while the \( y \) (\( \alpha \) ) axis of well \( A \) is parallel to the \( y \) (\( x \) ) axis of well \( B \) when the jawing angle is 0°.) If both the excited state of the donor and the acceptor state can be approximately described as the same single-configuration states (see Appendix B), a slow forward rate would be expected due to an electronic orientation effect, even when the jawing angle is small and the porphyrins are face-to-face, as in the Chang compounds. With \( \Phi_s(\phi) = \cos m\phi \) in each well, both the \((5,\pi) \rightarrow (5,\pi)\) and \((5,\pi) \rightarrow (4,\pi)\) transfers would yield zeros for \( H_{BA} \). Calculations are not shown since both transfers yield \( H_{BA} = 0 \). This feature is again an orbital shape effect peculiar to this class of orientations and results from the orthogonality of

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(57) In the event that the reaction is nonadiabatic in both the forward and reverse directions, this avoided crossing in the inverted region will not cause any greater decrease in the rate than does the nonadiabaticity (i.e., the small splitting) in the normal region. In this case the major difference between forward and reverse rates would be due to the inverted effect rather than the avoided crossing.
the $\Phi_m(\psi)'s$. In the rotated configuration, $\phi_B = \frac{1}{2} \pi - \phi_A$, so that $\cos 5\phi_B = \sin 5\phi_A$, which is itself orthogonal to $\cos 5\phi_A$. For nonzero jwelling angles $H_{BA}$ is still predicted to be zero for both forward and back transfer within the present model.

To obtain a zero forward transfer matrix element for this class of orientations, the adjustment of Overfield et al.\textsuperscript{[11,12]} within the present model, it is necessary that the initially excited donor state be well characterized by a single-excitation wave function.

To the extent that this is not true, the present model would predict a nonzero forward transfer matrix element. The high intensity of the Q bands in the spectra of ref 11 leads one to expect a largely single-configuration excited state for the compounds studied there on the basis of the four-orbital model.\textsuperscript{30} Therefore, the results of Overfield et al.\textsuperscript{[11,12]} regarding the slow forward rate of transfer as compared to the compounds of Chang,\textsuperscript{7} may arise from an orbital orientation effect.

At present there are apparently no experimental results for systems having the orientations of Figures 5c and 5d, corresponding to the results presented in Figures 9–11. In the results of Figure 9, in which an edge-to-edge orientation was considered, it is seen that the shape of the $H_{BA}$ vs. $\Gamma$ plot is relatively unaffected by an increase in separation distance, although $H_{BA}$ itself rapidly decreases with this distance.

In the calculations presented in Figures 10 and 11, $H_{BA}$ is examined at fixed edge-to-edge separations as a function of the angle $\Theta$. The plots for both the $(5, \pi)$ and $(5, \pi)$ and $(4, \pi)$ transfers exhibit several zeros and maxima between $\Theta = 0^\circ$ and $\Theta = 90^\circ$. At $\Theta = 0^\circ$ the forward transfer $H_{BA}$ shows a relative maximum while the back transfer $H_{BA}$ is zero, the explanation being that given for the $\delta = 0$ results of Figures 6 and 7. The orbital shape effect responsible for the difference in forward and back transfer $H_{BA}$'s for a face-to-face orientation ($\Theta = 0^\circ$) is not operative in the edge-to-edge configuration ($\Theta = 90^\circ$). The results illustrate the sensitivity of $H_{BA}$ not only to the states involved in the transfer but also to the molecular orientation. The change in relative size of the several maxima in Figures 10 and 11 as a function of separation distance can be qualitatively understood on the basis of orbital shape arguments similar to those given earlier.\textsuperscript{87}

The orientations examined in Figures 12 and 13 are of interest to discussions of forward and reverse electron transfer in bacterial photosystems. For the approximate experimental geometry\textsuperscript{6} ($\Delta = \Delta = 70^\circ$) the back transfer matrix element is more or less comparable in magnitude to that for forward transfer. Figure 13 indicates that no appreciable lowering of $H_{BA}$ for back transfer relative to that for forward transfer can be obtained by the change in orientation examined there. In the present model, therefore, there is no indication of electronic effects which would control the relative rates of forward and back electron transfer between the initial donor and the nearest BChl $b$ monomer in bacterial photosynthesis.

Other factors may then be responsible for the control of the relative forward and reverse rates in this photosynthetic system as discussed earlier, for instance the inverted effect.\textsuperscript{55} These back electron transfers are excellent candidates for obtaining an inverted effect, due to the expected low reorganization energies\textsuperscript{7–19} and the high driving forces. Also, the exact role of the BChl $b$ in transfer to the adjacent BPh has not been unambiguously established. It is possible that the BPh is involved in the initial electron transfer, in which case the orientations of all three compounds should be considered, perhaps within a superexchange mechanism,\textsuperscript{20,21,58,59} with the intermediate BChl $b$.

In general, the present results exhibit several maxima and zeros in $H_{BA}$ as a function of the variation of a given orientational parameter. To the extent that the $\pi$-orbitals of the actual systems have shapes similar to the model wave functions used here and to the extent that many-electron effects can be neglected, qualitative agreement of more elaborate theoretical studies with the present results can be expected. However, some deviations from the actual positions of the maxima would not be unexpected.

We have noted that the present one-electron model predicts a large difference in forward and back transfer $H_{BA}$'s in the face-to-face configuration. It is useful to inquire how model dependent this is. In $D_{2h}$ and $D_{2}$ porphyrins, the HOMO and LUMO are predicted to belong to a different irreducible representation of the molecular point group,\textsuperscript{88,42,43} and thus will be orthogonal in the face-to-face configuration not only for the present wave functions but also for more detailed ones. If the actual many-electron potential can be approximated by a reasonably smooth one, then it would be expected that back transfer would again be predicted to be slow by using these more detailed wave functions in the face-to-face configuration. Deviations from $D_{2h}$ or $D_{2}$ symmetry do not appear to affect the general shape of the HOMO or LUMO orbitals in ab initio calculations,\textsuperscript{41–43} so it is reasonable to expect that this particular prediction for the face-to-face configuration is not highly model dependent.

It might appear from eq 8 that $H_{BA}$ depends on the experimental "tail" of $\Psi_A$ but not on that of $\Psi_B$. However, it actually depends on the latter, albeit indirectly, via the $V_{s}$ appearing in that equation. $H_{BA}$ can also be written equivalently as $-V_{s}^{A} \langle \Psi_{A} | \Psi_{B} \rangle \lambda$, for the present case, where $\langle \Psi_{A} | H_{BA} | \Psi_{B} \rangle = E = \langle \Psi_{A} | H_{BA} | \Psi_{A} \rangle$.

The medium between two reactants is sometimes ordered and sometimes disordered. While medium effects can be considered as included in the present model via the adjustment of $E$ to yield a given value of $\beta$, the detailed effects of such environments on modifying, in a superexchange mechanism, the broad picture of the relative orientation effects described in this paper have not been examined. The general effects are expected to persist nevertheless. Clearly, experimental results when they become available will be particularly helpful in defining the practical utility of the present model and its predictions.

V. Conclusions

Calculations were reported for mutual orientation effects in electron transfers in diporphyrin systems. The donor and acceptor states were modeled by using one-electron eigenfunctions of oblate-spheroidal wells. A variety of orientations were examined, and it was shown that the thermal matrix element for electron transfer is a highly sensitive function of orientation and the orbits involved in the transfer. As more experimental results on such systems become available, the validity of the present predictions can be assessed.

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Appendix A: Evaluation of $H_{BA}$ as a Surface Integral

A simplified method for evaluating $H_{BA}$ is presented here, based on a method introduced by Bardeen.\textsuperscript{60} The method is applicable to all geometries of nonoverlapping wells.

In the present model, $H_{BA}$, the main contribution to the thermal matrix element for electron transfer may be written as a volume integral over well B. That is

$$H_{BA} = -V_{s}^{B} \int_{\text{well B}} \Psi_{A}^{*} \Psi_{B} \, d\tau$$

where $d\tau$ signifies a three-dimensional integral. In well B, $-V_{s}^{B} \Psi_{B}^{*}$ equals $(E - T) \Psi_{B}^{*}$ where $T$ is the one-electron kinetic energy operator. (We write $E$ rather than $E_{B}$, since we consider...
The number in parentheses denotes the electron in the given orbital. The symbol $\mathcal{A}$ is the antisymmetrizer operator which can be included by allowing for configuration interaction. Within particle interchange. The effects of electron-electron interactions guarantees that the wave functions are antisymmetric under single excitations, for transitions polarized in the $x$ direction, are by the four-orbital model it is assumed that only the above two primitive excitations contribute significantly to the lowest excited singlet state. The secular equation which then determines the multiconfiguration excited state is

$$
\begin{bmatrix}
E_1 & \frac{H_{12}}{H_{11}} \\
\frac{H_{12}}{H_{11}} & E_2
\end{bmatrix}
\begin{bmatrix}
C_1 \\
C_2
\end{bmatrix}
= E
\begin{bmatrix}
C_1 \\
C_2
\end{bmatrix}
$$


$$
E_1 \text{ and } E_2 \text{ are the energies of the primitive excitation functions in eq B1, and } H_{12} = (H_{21}) \text{ is a two-electron interaction term. The solutions of eq B2 are linear combinations of } \psi_1 \text{ and } \psi_2 \text{ which are then taken to model the states observed in the } x \text{-polarized } Q \text{ and } B \text{ bands of the porphyrin. In the present model } E_1 = E_2 \text{ so the states obtained lead to } C_1 = \pm C_2. \text{ However, upon examination of the asymmetric systems the spectra indicate that } E_1 \neq E_2 \text{ in general.}$$

The thermal matrix element $H_{BA}$ is

$$
H_{BA} = \langle \Psi_{\text{donor}} | V_{\text{acceptor}} | \Psi_{\text{acceptor}} \rangle
$$

It is assumed that $\Psi_{\text{donor}}$, the wave function before electron transfer, corresponding to the first excited singlet state, is

$$
\Psi_{\text{donor}} = \mathcal{A} \left( \cos 4\varphi(1) \cos 5\varphi(2) - \sin 4\varphi(1) \sin 5\varphi(2) \right) \Phi
$$

Substituting in eq B3, $H_{BA}$ becomes

$$
H_{BA} = \mathcal{A} \left( \cos 4\varphi(1) \cos 5\varphi(2) - \sin 4\varphi(1) \sin 5\varphi(2) \right) \Phi |V_{\text{acceptor}}| \Psi_{\text{acceptor}}
$$

Since the wave functions here are two-electron wave functions, $V_{\text{acceptor}}$ is of the form $V_{\text{pH1}}(A) + V_{\text{pH2}}(B)$. In choosing the wave function for the final state, $\Psi_{\text{acceptor}}$, it is assumed that the electronic state on the acceptor can be represented by a single configuration, say $5\varphi_B$. Similarly, the oxidized donor is represented by a single-configuration state, either $4\varphi_D$ or $5\varphi_D$. Then the possible $\Psi_{\text{acceptor}}$ states are (again neglecting all doubly occupied orbitals)

$$
\Psi_{\text{acceptor}} = \left\{ \begin{array}{c}
\mathcal{A} \left( \cos 4\varphi(1) \cos 5\varphi(2) \right) \\
\mathcal{A} \left( \sin 4\varphi(1) \cos 5\varphi(2) \right)
\end{array} \right\}
$$

If one chooses $\Psi_{\text{acceptor}} = \mathcal{A} \left( \cos 4\varphi(1) \cos 5\varphi(2) \right) \Phi$ and when $(\sin 5\varphi(2) \cos 5\varphi(2)) = 0$, as is the case for all orientations in the present article, $H_{BA}$ becomes

$$
H_{BA} = \frac{1}{21/2} \left( \mathcal{A} \cos 4\varphi(1) \cos 5\varphi(2) \times |V_{\text{acceptor}}| \mathcal{A} \cos 4\varphi(1) \cos 5\varphi(2) \right)
$$

In obtaining eq B7, we have performed the integrations over the spin variables. By expansion of eq B7, the dominant terms sum to yield (dropping the dummy indices)

$$
H_{BA} = \frac{1}{21/2} \left( \cos 4\varphi(1) \cos 5\varphi(2) \right)
$$

The quantity in eq B8 is just $1/21/2$ times the one-electron matrix elements given in the present paper. Therefore, the one-electron matrix elements given in the text are sufficient to discuss the orientation dependence of electron transfers involving such multiconfiguration states.

In eq B4-B8 the donor state is multiconfigurational and the acceptor state is single-configurational (the former justified as in ref 38 and 39 and the latter justified as in ref 45 and 46). The reason why the $|H_{BA}|^2$ obtained from eq B8 differs by a factor of 2 from the single-configuration value used in the text is that in the former the donor wave function contains both cosine and sine components, one of which by symmetry does not contribute to $H_{BA}$ when the acceptor state is a single configuration.