

Energetics of Quinone-Dependent Electron and Proton Transfers in *Rhodobacter sphaeroides* Photosynthetic Reaction Centers

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ABSTRACT: Proteins bind redox cofactors, modifying their electrochemistry and affinity by specific interactions of the binding site with each cofactor redox state. Photosynthetic reaction centers from *Rhodobacter sphaeroides* have three ubiquinone-binding sites, Q_A , and proximal and distal Q_B sites. Ubiquinones, which can be doubly reduced and bind 2 protons, have 9 redox states. However, only Q and Q^- are seen in the Q_A site and Q , Q^- , and QH_2 in the proximal Q_B site. The distal Q_B function is uncertain. Multiple conformation continuum electrostatics (MCCE) was used to compare the ubiquinone electrochemical midpoints (E_m) and pK_a values at these three sites. At pH 7, the Q_A/Q_A^- E_m is -40 mV and proximal Q_B/Q_B^- -10 mV in agreement with the experimental values (assuming a solution ubiquinone E_m of -145 mV). Q_B reduction requires changes in nearby residue protonation and SerL223 reorientation. The distal Q_B/Q_B^- E_m is a much more unfavorable -260 mV. Q_A and proximal Q_B sites generally stabilize species with a -1 charge, while the distal Q_B site prefers binding neutral species. In each site, the dianion is destabilized because favorable interactions with the residues and backbone increase with charge (q), while the unfavorable loss of solvation energy increases with q^2 . Therefore, proton binding before a second reduction, forming QH and then QH^- , is always preferred to forming the dianion (Q^{2-}). The final product QH_2 is higher in energy at the proximal Q_B site than in solution; therefore, it binds poorly, favoring release. In contrast, QH_2 binds more tightly than Q at the distal Q_B site.

Quinones are found in many transmembrane proteins that coupled electron and proton transfer reactions (1, 2), including photosynthetic reaction centers, PSII (3), PSI (4), cytochrome bc_1 (5), and b_6f complexes (6–8), and in the quinol:fumarate (9) and succinate:quinone (10) oxidoreductases. With their ability to bind 2 protons and 2 electrons, quinones have 9 redox states (Figure 1) (11). In aqueous solution at physiological pH, these quinones go from being fully oxidized (Q) to fully reduced (QH_2) in a concerted two electron and two proton reaction (11). This electrochemistry is modified within individual proteins; therefore, reactions can proceed via single electron transfers so reduction steps do not need to be coupled to proton uptake.

The quinone-dependent electron and proton transfers in bacterial reaction centers (RCs) of *Rhodobacter sphaeroides* have been well-studied (1, 2, 12), providing a wealth of information that can be subjected to computational analysis (13). This theoretical analysis enhances our understanding of the protein while providing good tests of evolving computational methods. RCs consist of three protein subunits, L, M, and H, and 10 cofactors including 2 ubiquinones. The intraprotein electron transfer occurs along the L branch from the excited singlet bacteriochlorophyll dimer (P), through a bacteriochlorophyll monomer (B) and bacteriopheophytin monomer (H) to the primary quinone $Q_A \approx 25$ Å away in 150 ps. Q_A then reduces the secondary quinone Q_B 15 Å away (edge to edge) in 100 μs (14, 15). Cytochrome c reduces P^+ , yielding $PQ_A Q_B^-$. After a second excitation of

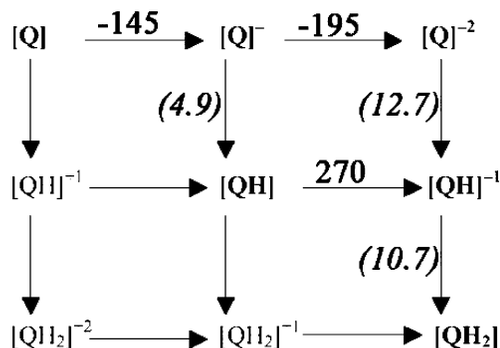


FIGURE 1: Ubiquinone redox and protonation states. States in bold are intermediates considered in coupled electron (horizontal) and proton (vertical) transfer reactions in RCs. The sources for the $E_{m,sol}$ and $pK_{a,sol}$ (in parentheses) at pH 7 are given in Table 1. The ΔG for proton transfer (vertical lines) can be obtained from $\Delta G = 59.3 \cdot (pH - pK_a)$ (eq 9).

P, $P^+Q_A^-Q_B^-$ is formed. There is good evidence that Q_B binds a proton (forming Q_BH) prior to its second reduction (16). The second reduction of Q_B then forms Q_BH^- , which binds a second proton, yielding the dihydroquinone. Overall, two cycles of light-driven reactions oxidize two cytochromes c in the periplasm, delivering these two electrons along with two protons from the cytoplasm to Q_B . The reduced dihydroquinone is released to the cell membrane, and a new oxidized quinone is bound (17). In the cell, electron transfer from the dihydroquinone back to cytochrome c occurs in the cytochrome bc_1 complex, which adds to the transmembrane proton gradient by releasing protons into the periplasm.

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The protein modifies the ubiquinone electrochemistry to differentiate Q_A and Q_B . In the Q_A site, only the oxidized Q_A and anionic semiquinone Q_A^- are found. Q_A does not dissociate from the protein. Q_B displays more complex redox chemistry, serving as the two-electron gate (18). Thus, two single electron transfers via Q_A form the doubly reduced Q_B for release into the membrane (12). RCs can be found with Q_B in three relatively stable redox states: unreduced quinone (Q), anionic semiquinone (Q^-), and fully reduced and protonated dihydroquinone (QH_2). The anionic semiquinone is tightly bound to the protein, while the Q and QH_2 freely exchange with the quinone pool in the membrane (19). The pathway for the second reduction indicates that of the two possible intermediates Q_BH is easier to form than Q_B^{-2} ; therefore, proton binding occurs prior to electron transfer. Thus, of the nine possible redox states for Q_B , five are found on the reaction pathway (Figure 1). In addition, in the reaction scheme, it is necessary that the redox potentials of Q_B for the first and second reduction are maintained so that Q_A^- can reduce both Q_B and Q_BH . Lastly, there are two binding sites for Q_B , distal and proximal, seen in the crystal structures (20, 21). Kinetic measurements find no evidence for quinone reduction in the outer, distal site (22–26), although this has been proposed by simulation (27). A comparison of the thermodynamics of quinone reduction in the two sites can help to determine what redox states are accessible in the distal site.

The different *in situ* thermodynamic properties of Q_A and Q_B can be attributed to different local environments. Q_B is surrounded by a cluster of polar residues, acids, and waters, including SerL223, AspL210, GluL212, AspL213, GluH173, AsnM44, and AspM17. These residues have been identified as important for electron and proton transfer by both mutation experiments (28–36) and simulations (37, 38). These residues rearrange to stabilize the anionic semiquinone and provide a pathway for proton delivery prior to and following the second reduction of Q_B . This flexible polar region of the protein maintains the appropriate energies of the different Q_B redox states. In contrast, there are fewer ionization shifts and no proton pathway associated with reduction of Q_A .

With the wealth of experimental information about the redox chemistry in wild-type and mutant protein, RCs provide an excellent system to test simulation techniques. Molecular dynamics (MD) has been used to study Q_B movements (39), conformational gating (40), and changes in protonation states of amino acids GluL212 and AspL213 on the first electron transfer (41). Continuum electrostatics (CE) simulations provide rapid estimates of long-range electrostatic interactions by averaging many motions into the dielectric constant (13, 42). The first electron transfer from Q_A^- to Q_B has been studied by various CE methods using both *Rhodospseudomonas viridis* (43, 44) and *Rb. sphaeroides* RC structures (37, 38, 45–47). The energetics of the second Q_B reduction has also been calculated by Knapp and co-workers with a CE model in *Rps. viridis* (44) and *Rb. sphaeroides* (46) RCs.

Standard CE techniques provide two types of response to changes in charge caused by reactions such as quinone reduction (13, 42, 48). One is a homogeneous dielectric response defined by the dielectric constant used in the calculation. Values from 4 to 20 have been used with the higher values averaging more conformational changes in the

protein. The other is the changes in protonation of surrounding residues. These proton shifts coupled to electron transfer are analyzed using standard methods to calculate residue pK_a values (48–50). Here, no explicit conformational changes are allowed. Multiple conformation continuum electrostatics (MCCE) calculations add side-chain heavy atom and hydroxyl conformers to improve the flexibility of the CE model (51–54). This method can follow linked conformational and ionization state changes during the reduction of quinone. The work presented here uses MCCE to analyze the energies of the redox states in the Q_A and distal and proximal Q_B sites to determine how the protein stabilizes the appropriate states in each binding site.

MATERIALS AND METHODS

MCCE Methods. MCCE (13, 37, 53, 54) calculates the equilibrium distribution of conformation and ionization states of protein side chains, buried waters, ions, and ligands at a defined solution pH and E_h . Preselected choices for atomic coordinates and ionization states are used. Side-chain conformers are added providing alternative positions of hydroxyl protons of Ser, Tyr, Thr, and water, His tautomers, and the different ionization states of the acids Asp, Glu, and Tyr and the bases Arg, Lys, and His. There are at least two Q_B -binding sites (20), characterized in a comparison of protein frozen in the light (1AIG) and dark (1AIJ) (21, 55). The protein structure (1AIJ) with a resolution of 2.2 Å used in this calculation has the quinone in the distal position. The proximal Q_B position of 1AIG was added into the 1AIJ structure by superimposing the Q_B ring, non-heme iron, and the backbone atoms of GluL212 and AspL213. Conformers of protonated quinone species are generated with 2 different hydrogen positions for each carbonyl oxygen in the plane of the quinone ring with an angle of 60° between them (Figure 5). This generates 4 neutral semiquinone (QH) and 4 dihydroquinone (QH_2) conformers.

Lipid and detergent molecules are removed from the structure. The 110 waters with <10 Å exposed surface area defined by SURFV (56, 57) are included. Conflicts between waters at nearby positions in a given site are avoided by providing each water with a conformer that does not interact with the protein. Thus, each water molecule is allowed to leave the protein during Monte Carlo sampling. Hartree–Fock partial charges are used for each ubiquinone redox state (44). A +2 charge is placed on the non-heme iron between Q_A and Q_B . The partial charges for all other cofactors are from ref 58. PARSE partial charges and radii (59) are used for all other groups. For calculations involving Q_A or proximal Q_B , no quinone conformers are allowed in the distal Q_B site (Q_{BD}). For analysis of Q_{BD} , the proximal Q_B site is kept unoccupied. For states where the quinone is protonated, conformers with different proton positions are allowed and their occupancy is summed. Unless otherwise noted, Q_A is fixed in the oxidized state for all Q_B titrations, and likewise, the proximal Q_B is fixed in the quinone state and Q_{BD} is deleted, for Q_A titrations.

Look-up tables are calculated for electrostatic and non-electrostatic conformer self- and conformer–conformer pairwise interactions. The electrostatic pairwise interactions and reaction field energies are calculated using the finite-difference technique to solve the Poisson–Boltzmann equa-

tion with DelPhi (57, 60, 61). Three focusing runs are done, giving a final resolution of 1.2 grids/Å (62). The protein is surrounded by water. The protein dielectric constant is 4, while 80 is used for the solvent with a salt concentration of 150 mM. The Lennard–Jones interactions are calculated using A and B parameters previously described (37). The possible microstates of the system are subjected to Monte Carlo sampling. A microstate is made up of one conformer for each residue, cofactor, and water. The energy of microstate n (ΔG^n)¹ is the sum of the electrostatic and nonelectrostatic energies (63) (eq 1), where $k_b T$ is 0.59 kcal/

$$\Delta G^n = \sum_{i=1}^M \delta_x(i) \{ \gamma(i) [2.3k_b T(\text{pH} - \text{p}K_{\text{sol},i}) - nF(E_h - E_{\text{m},\text{sol},i})] + (\Delta\Delta G_{\text{rxn},i} + \Delta G_{\text{pol},i} + \Delta G_{\text{nonel},i}) \} + \sum_{i=1}^M \delta_x(i) \sum_{j=i+1}^M \delta_x(j) [\Delta G_{ij}^{\text{el}} + \Delta G_{ij}^{\text{nonel}}] \quad (1)$$

mol (25.8 meV), M is the number of conformers, $\delta_x(i)$ is 1 for conformers that are present in the state and 0 for all others, $\gamma(i)$ is 1 for bases, -1 for acids, and 0 for polar groups and waters, n is the number of electrons gained or lost in redox reactions, F is the Faraday constant, $\text{p}K_{\text{sol},i}$ is the $\text{p}K_a$ of the i th group in solution, $E_{\text{m},\text{sol},i}$ is the midpoint potential of the i th cofactor in solution, $\Delta\Delta G_{\text{rxn},i}$ is the difference between the conformer reaction field energy in solution and the protein (the desolvation energy), $\Delta G_{\text{pol},i}$ is the pairwise electrostatic interaction of a conformer with the backbone and with side chains that have no conformational degrees of freedom, $\Delta G_{\text{nonel},i}$ is the Lennard–Jones interactions with the backbone and with all side chains with no degrees of freedom plus the torsion angle energy, and $\Delta G_{ij}^{\text{el}}$ and $\Delta G_{ij}^{\text{nonel}}$ are the electrostatic and Lennard–Jones pairwise interactions between each conformer in the microstate. The limits on the summation of the interconformer terms ensure that each interaction is counted once. Monte Carlo sampling establishes the Boltzmann distribution of different conformers of each residue at 25 °C, providing the probabilities of the quinone reactant and product states at a given solution redox potential (E_h) and pH. Multiflip (64) between closely coupled residues is implemented (65). The SOFT function is not used (54). A total of 40 million steps of Monte Carlo sampling leads to convergence for the system with 449 residues, including waters, with conformational degrees of freedom. There are 3833 conformers to be sampled. For each reported E_m , three Monte Carlo runs were made and the results were averaged. The average uncertainty in the conformer occupancy is ± 0.04 . The uncertainty of calculated E_m values is ± 4 mV.

Redox Potentials and $\text{p}K_a$ Values of Quinones in Solution. MCCE calculates the shift in free energy of a site redox or protonation reaction when it is moved from water into the protein, providing the changes in the electrochemical midpoint, E_m , or $\text{p}K_a$. Given a reference, $E_{\text{m},\text{sol}}$ or $\text{p}K_{\text{a},\text{sol}}$ for the cofactor in solution, measurable E_m and $\text{p}K_a$ values in the

protein can be calculated (63). The redox midpoint potential of UQ/UQ^- and $\text{UQ}^-/\text{UQ}^{2-}$ cannot be measured in aqueous solution at pH 7 because the $E_{\text{m},\text{sol}}$ for quinone reduction to the semiquinone is lower than that for formation of the fully protonated dihydroquinone (QH_2). Thus, in water at physiological pH, UQ is reduced in an $n = 2$ reaction to QH_2 (11, 66). Measurements of the one electron reactions have been made for a small number of quinones in water (67). In the absence of protons, the semiquinone is stable; therefore, E_m measurements are far more straightforward. E_m values for Q/Q^- have been determined in the aprotic solvent dimethylformamide (DMF) for a large number of quinones (68).

Several of the aqueous $E_{\text{m},\text{sol}}$ for ubiquinones used here were estimated by comparison with the E_m values of tri- and tetramethyl benzoquinones. UQ, a 2,3-dimethoxy-5-methyl-6-isoprenyl benzoquinone, is considered to behave more like trimethyl- than tetramethyl benzoquinone because the 2,3-methoxys cannot both lie in the plane of the ring. Therefore, while two adjacent methyl groups can be electron-donating into the ring, two methoxy group cannot be (68). In DMF, trimethyl benzoquinone has an E_m for the half-reaction $\text{Q} + e^- \rightarrow \text{Q}^-$ that is 30 mV more negative than UQ, while that of the tetramethyl compound is 150 mV lower. The E_m for the trimethyl semiquinone is -170 mV in water (67). An $E_{\text{m},\text{sol}}$ for UQ of -145 mV, 30 mV more positive than trimethyl benzoquinone, will be used here.

The E_m (Q/Q^-) for ubiquinone in DMF was measured to be -360 mV (68). The value of -145 mV represents a shift of 215 mV moving from water to DMF as found for many small quinones (69). A pure CE analysis of the change in the Born reaction field energy on moving the Q to Q^- reaction from DMF ($\epsilon = 37$) to water ($\epsilon = 80$) only predicts a shift stabilizing the anionic semiquinone by ≈ 40 meV, shifting the $E_{\text{m},\text{sol}}$ to -320 mV (44). Thus, more specific interactions of water with the semiquinone not considered in the CE analysis are assumed to shift the E_m up to -145 .

The aqueous $\text{p}K_{\text{a},\text{sol}}$ of semiubiquinone (UQ^-/UQH) was estimated to be 4.9 (67). Higher values of 6.5 are found in 80% ethanol/water (w/w) (70, 71). The $\text{p}K_a$ (QH^-/QH_2) of UQ was measured as 13.3 in 80% ethanol/water (w/w) (70). Using one suggested correction of -1.6 pH units moving from 80% ethanol to water places the ubiquinone $\text{p}K_{\text{a},\text{sol}}$ at ≈ 11.7 , a high value given the electronegativity of the methoxy substituents. Rather, the aqueous, high pH trimethyl benzoquinone $\text{p}K_{\text{a},\text{sol}}$ of 10.7 will be used here (72). The gap between $\text{p}K_{\text{a},\text{sol}}$ (QH^-/QH_2) and $\text{p}K_{\text{a},\text{sol}}$ ($\text{Q}^{2-}/\text{QH}^-$) is rather constant at ~ 1.5 – 2 units (71, 72). This places the $\text{p}K_{\text{a},\text{sol}}$ ($\text{Q}^{2-}/\text{QH}^-$) at 12.7 (69). Combining these $\text{p}K_a$ values and the $E_{\text{m},7}$ for $\text{Q}^- + e^- + 2\text{H}^+ \rightarrow \text{QH}_2$ of -360 mV (69) provides enough information to derive an $E_{\text{m},\text{sol}}$ for $\text{Q}^- + e^- \rightarrow \text{Q}^{2-}$ of -195 mV as well as the other values in Table 1. The E_m (Q^-/Q^{2-}) for UQ_0 in DMF is -1080 mV (68, 69).

Calculation of the Free Energy of Quinone Reduction Reactions. The free energy of quinone reduction or protonation reactions in RCs, ΔG , at a given pH and E_h is

$$\Delta G = 2.3mk_b T(\text{pH} - \text{p}K_a) - nF(E_h - E_m) = -2.3k_b T \log \frac{[P]}{[R]} \quad (2)$$

where F is the Faraday constant, m and n are the number of

¹ Abbreviations: All free-energy terms ΔG and electrochemical midpoints (E_m) refer to standard conditions at pH 7 ($\Delta G^{\circ'}$ and $E_{\text{m},7}^{\circ'}$). The energy to change a $\text{p}K_a$ by 1 pH unit at 20 °C is 1.36 kcal/mol = 59.3 meV. $\text{p}K_a'$ is the $\text{p}K_a$ calculated from the free energy of protonation at pH 7.

Table 1: Reference E_m and pK_a Values for Ubiquinone 10

half-reactions	$pK_{a,sol}$	$E_{m,sol,7}^{\circ}$ (mV)	ΔG_{sol}° (meV)
$Q + e^- \rightarrow Q^-$		-145 ^b	145
$Q + 2e^- \rightarrow Q^{-2}$		-170 ^b	340
$Q^- + e^- \rightarrow Q^{-2}$		-195 ^c	195
$QH + e^- \rightarrow QH^-$		270 ^b	-270
$Q^- + H^+ \rightarrow QH$	4.9 ^d		124
$Q^{-2} + H^+ \rightarrow QH^-$	12.7 ^e		-338
$QH^- + H^+ \rightarrow QH_2$	10.7 ^f		-218
$Q + e^- + H^+ \rightarrow QH$		-269	269
$Q + 2e^- + H^+ \rightarrow QH^-$		1	-1
$Q + 2e^- + 2H^+ \rightarrow QH_2$		220	-220
$Q^- + e^- + H^+ \rightarrow QH^-$		146	-146
$QH + e^- + H^+ \rightarrow QH_2$		488	-488
$Q^- + e^- + 2H^+ \rightarrow QH_2$		360	-360

^a ΔG_{sol} is $2.3mk_bT(pH - pK_{a,sol}) - nF(E_h - E_{m,sol})$ (eq 4), and values are given at $E_h = 0$ and $pH 7$, where m is the number of protons and n is the number of electrons. ^b By analogy with the E_m of trimethyl benzoquinone (67). ^c Derived from $E_{m,sol}(Q/Q^-)$ and $E_{m,sol}(Q^-/Q^{-2})$. ^d Derived from the $pK_{a,sol}$ for Q^{-2}/QH^- and the $E_{m,sol}$ for Q^-/Q^{-2} . ^e Derived from $pK_a(Q^{-2}/QH^-)$ being ≈ 2 units higher than $pK_a(QH^-/QH_2)$ (69, 72). Other values are derived as appropriate sums and differences walking around the thermodynamic box (Figure 1). See the Materials and Methods for a more complete derivation of $E_{m,sol}$ and $pK_{a,sol}$ values used here. ^f ref 67.

protons and electrons transferred, respectively, pK_a and E_m are values for quinone in the protein, and $[R]$ and $[P]$ are the occupancies of the reactant and product, respectively, calculated by Monte Carlo sampling given the microstate energies in eq 1. To determine the *in situ* ΔG for each reaction, only conformers for given reactant and product redox states are sampled. For example, to determine ΔG for the reaction $Q_B H^- + e^- + H^+ \rightarrow Q_B H_2$, only conformers of $Q_B H^-$ and $Q_B H_2$ are allowed. Q_A would be fixed in its oxidized form, and no conformers of Q_{BD} would be allowed in any sampled microstate. There are four conformers with different locations for the protons on $Q_B H^-$ and $Q_B H_2$ (see Figure 5 below). Their accepted occupancies are summed. The reaction free energy is divided into two terms

$$\Delta G = \Delta G_{sol} + \Delta \Delta G_{protein} \quad (3)$$

ΔG_{sol} is the reference free energy in solution at the given pH and E_h . This includes the contributions of the intrinsic quinone proton or electron affinity and the ability of the solvent to donate protons and electrons given the pH and E_h . Thus

$$\Delta G_{sol} = 2.3mk_bT(pH - pK_{a,sol}) - nF(E_h - E_{m,sol}) \quad (4)$$

$pK_{a,sol}$ and $E_{m,sol}$ are the quinone reference values in water (Figure 1 and Table 1). $\Delta \Delta G_{protein}$ is the shift in the energy of reduction or protonation as the quinone is moved from water into the protein (13, 63). It is calculated from

$$\Delta \Delta G_{protein} = -2.3k_bT \log \frac{[P]}{[R]} - \Delta G_{sol} \quad (5)$$

$\Delta \Delta G_{protein}$ is due to the changes in quinone pK_a ($\Delta pK_{a,protein}$) and/or E_m ($\Delta E_{m,protein}$). Thus

$$\Delta \Delta G_{protein} = -2.3mk_bT \Delta pK_{a,protein} + nF \Delta E_{m,protein} \quad (6)$$

As the E_h titrations are carried out at pH 7, $\Delta \Delta G_{protein}$ is $nF \Delta E_{m,7}$. The E_m and pK_a values calculated for the quinone

in the protein are each viewed as the protein shifting the solution quinone behavior. Thus

$$E_m = E_{m,sol} + \Delta E_{m,protein} \quad (7)$$

$$pK_a = pK_{a,sol} + \Delta pK_{a,protein} \quad (8)$$

For the proton independent half-reactions ($Q + e^- \rightarrow Q^-$, $Q^- + e^- \rightarrow Q^{-2}$, and $QH + e^- \rightarrow QH^-$) $m = 0$ (eq 6); therefore, $\Delta \Delta G_{protein}$ is simply related to the shift in E_m in the protein, $\Delta E_{m,protein} = -\Delta \Delta G_{protein}/nF$ (63). When determining $\Delta \Delta G_{protein}$ for the proton-coupled electron transfers (e.g., $Q + e^- + H^+ \rightarrow QH$), $\Delta \Delta G_{protein}$ includes contributions from both $\Delta pK_{a,protein}$ and $\Delta E_{m,protein}$.

Quinone pK_a values can be determined by Monte Carlo sampling as a function of pH (54). However, this will not yield the correct free energy of protonation at pH 7 because the protein ionization and conformation are pH-dependent. The free energy of proton-transfer reactions at pH 7 was obtained from the difference between the energy of the electron transfer and the coupled electron- and proton-transfer reaction closing the thermodynamic cycle (Figures 1 and 3) (73). For example, the standard free energy of the protonation reaction $Q_B^- + H^+ \rightarrow Q_B H$ at pH 7 is the difference between the ΔG of $Q_B + e^- + H^+ \rightarrow Q_B H$ and $Q_B + e^- \rightarrow Q_B^-$. A pK_a' , the effective pK_a calculated at pH 7, is then obtained from the pH where this ΔG for the protein would be zero if the protein remained as it is at pH 7

$$pK_a' = 7 - \Delta G/RT \quad (9)$$

The half-reaction E_m and pK_a values were obtained for the relevant quinone reactions in the three binding sites. The energy ΔG of a particular electron-transfer reaction such as from $Q_A^- Q_B$ to $Q_A Q_B^-$ is then the difference of the redox potentials $E_m(Q_A/Q_A^-)$ and $E_m(Q_B/Q_B^-)$.

Factors Contributing to $\Delta G_{protein}$. In the MCCE analysis, the free-energy change due to the protein (eq 6) can be broken down into

$$\Delta \Delta G_{protein} = (\Delta \Delta G_{rxn} + \Delta G_{pol} + \Delta G_{nonel}) + \Delta G_{res} \quad (10)$$

All terms are changes in the free energy of the reactant and product redox state of the quinone when it is moved from water to the protein. $\Delta \Delta G_{rxn}$, ΔG_{pol} , and ΔG_{nonel} are independent of the distribution of other conformers and are found in the energy look-up table for each conformer. ΔG_{res} , calculated with eqs 5 and 10, accounts for the changing interaction of the quinone with the protein as it undergoes the redox reaction, as well as the energy needed to keep the conformers and the rest of the protein in equilibrium with the quinone charge and protonation changes. Thus

$$\Delta G_{res} = \Delta G_{res}^{prot} + \Delta G_{prot \rightarrow prot^*}^{red} = \Delta G_{prot \rightarrow prot^*}^{ox} + \Delta G_{res}^{prot^*} \quad (11)$$

If, for example, the reactant is Q_B and the product is Q_B^- , ΔG_{res}^{prot} is the difference in interaction of the protein with Q_B and Q_B^- in the protein equilibrated around Q_B (prot), and $\Delta G_{prot \rightarrow prot^*}^{red}$ is the energy it takes to move the protein into the conformation equilibrated around Q_B^- when the Q_B is reduced (prot*). This is equivalent to reducing Q_B in the prot conformation and then relaxing the protein to the prot*

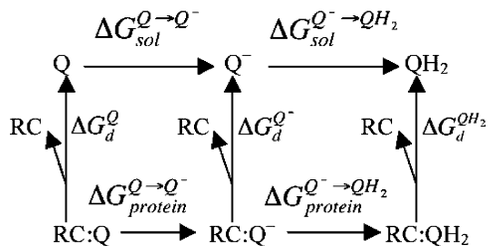


FIGURE 2: Relationship between the equilibrium dissociation energy (ΔG_d) and redox energies of the reaction in solution and in the protein. ΔG_d^Q , $\Delta G_d^{Q^-}$, and $\Delta G_d^{QH_2}$ are the dissociation free energies for Q, Q^- , and QH_2 , respectively.

conformation. The reduction could also happen by the alternate path, where first the protein is moved to the conformation equilibrated around Q_B^- , while keeping Q_B oxidized requiring $\Delta G_{\text{prot}^*}^{\text{ox}}$, and then the quinone is reduced in the pre-prepared protein, prot^* , ($\Delta G_{\text{res}}^{\text{prot}^*}$). $\Delta G_{\text{prot}^*}^{\text{ox}}$ is always unfavorable, while $\Delta G_{\text{prot}^*}^{\text{red}}$ is always favorable. $\Delta G_{\text{red}}^{\text{prot}^*}$ is always more favorable than $\Delta G_{\text{res}}^{\text{prot}}$ because the cost of rearranging into the protein equilibrated around the product has already been paid (see ref 63 for a more complete discussion).

Relative Affinity of Different Quinone Redox States. Each quinone redox state has a different affinity for each binding site required for RC function. To the degree that the protein stabilizes the reactant or product state in an electron- or proton-transfer process, it will bind that species more tightly (Figure 2) (74). For example, $\Delta\Delta G_{\text{protein}}$ for the $Q + e^- \rightarrow Q^-$ reaction provides a measurement of the relative affinities of Q and Q^- for the binding site

$$\Delta\Delta G_{\text{protein}} = \Delta G_d^Q - \Delta G_d^{Q^-} = \Delta G_{\text{protein}}^{Q \rightarrow Q^-} - \Delta G_{\text{sol}}^{Q \rightarrow Q^-} = 2.303RT(\log K_d^{Q^-} - \log K_d^Q) \quad (12)$$

ΔG_d is the dissociation energy for each species, and K_d is the derived dissociation constant for Q and Q^- .

RESULTS

Energy of Ubiquinone Reduction at Q_B Proximal, Q_B Distal, and Q_A Sites: First Reduction of Q_B in the Proximal Binding Site. The inner, proximal Q_B site is calculated to stabilize quinone reduction to the anionic semiquinone by -138 meV relative to the same reaction in water ($\Delta\Delta G_{\text{protein}}$) (values are given to the nearest millielectronvolt only as an aid for tracking the connections between the numbers). Using a solution reference E_m ($E_{m,\text{sol}}$) of -145 mV (Table 1), the E_m in the protein would be -7 mV (eq 7, Tables 2a and 5a) in good agreement with the experimentally determined value of ≈ 30 mV (75). $\Delta\Delta G_{\text{protein}}$ can be divided into the loss of stabilization of the charge by water when the cofactor moves from the solution ($\Delta\Delta G_{\text{rxn}}$), the electrostatic pairwise interactions with the protein backbone (ΔG_{pol}) and residues and ligands (ΔG_{res}), and the torsion and Lennard–Jones interactions (ΔG_{nonel}) (eq 10). ΔG_{res} includes the direct interactions between residues and the quinone as well as the energy required to change the residue conformation and ionization states so they remain in equilibrium in each quinone redox state (eq 11, Table 3).

For the first Q_B reduction, $\Delta\Delta G_{\text{rxn}}$ is ≈ 390 meV and ΔG_{pol} is ≈ -390 meV, with the backbone dipoles essentially

compensating for the removal from water (43, 76, 77) (Table 3). The primary contribution to ΔG_{pol} is from residues L210–247 (helix E), with smaller contributions from the stretches M136–162 and M232–288. Pairwise interactions with other side chains and ligands ultimately favor reduction by -140 meV (Table 2a). The $+2$ charge of the nearby non-heme iron provides a constant term favoring ionization of the proximal Q_B . However, other residues must undergo changes in ionization and conformation for Q_B^- to be stabilized.

Monte Carlo sampling in MCCE maintains the ionization states and position of surrounding residues in equilibrium with the quinone redox states. If the protein was fixed in the conformation and ionization equilibrated in the ground Q_B state, $\Delta G_{\text{res}}^{\text{prot}}$ is 86 meV (Table 3). Combining this with $\Delta\Delta G_{\text{rxn}}$, ΔG_{pol} , and ΔG_{nonel} , which are independent of the chosen conformers of other residues, yields $\Delta\Delta G_{\text{protein}}$ of 83 meV (eq 10). Adding the $E_{m,\text{sol}}$ of -145 mV yields an E_m for Q_B in the ground-state protein of -228 mV, indicating a very destabilized Q_B^- (eq 10, Table 3). This is consistent with earlier calculations (37) as well as the observation that in dark-adapted, frozen RCs Q_B cannot be reduced by Q_A (78, 79). It takes 183 meV to move the protein from the conformation equilibrated around Q_B to one equilibrated around Q_B^- without reducing the quinone ($\Delta G_{\text{prot}^*}^{\text{ox}}$) (Table 3). Reduction in the pre-prepared protein would be stabilized by $\Delta G_{\text{res}}^{\text{prot}^*}$ and would be very favorable, with a resultant E_m of 318 mV. The calculated reaction ΔG_{res} , -135 meV, is much less favorable (eq 11, Table 2a) because it includes the work to rearrange the protein from the reactant to product conformation.

Proton uptake to surrounding residues, in particular, GluL212 and AspL213, is the most important change required to stabilize Q_B^- . As found in earlier calculations (37), a net charge of -1 is maintained on these two acids and Q_B . In these simulations, when Q_B is oxidized, GluL212 is ionized and AspL213 is protonated (Appendix I in the Supporting Information). When Q_B is ionized, both acids are neutral. The orientation of the hydroxyl SerL223, which can offer a hydrogen bond to either AspL213 or Q_B also changes on reduction (Table 4). SerL223 hydroxyl positions that can donate a hydrogen bond to and accept a hydrogen bond from the neutral Q_B are accepted by Monte Carlo sampling (Table 4). When Q_B is reduced, SerL223 becomes a hydrogen-bond donor to the anionic semiquinone.

Protonation of the Anionic Semiquinone. QH and Q interact with the Q_B site similarly (Table 2a). The ionization of the acidic cluster of GluL212 and AspL213 are the same with $Q_B\text{H}$ or Q_B (Appendix I in the Supporting Information). The small differences in $\Delta\Delta G_{\text{rxn}}$ are due to differences in the charge distribution in the two quinone species (44). The differences in Lennard–Jones interactions (ΔG_{nonel}) and ΔG_{res} come from the interactions of the quinone proton with the protein. Three positions are acceptable with the majority of the quinones in Monte Carlo sampling donating a hydrogen bond to SerL223 (Table 4). The proximal Q_B site is designed to stabilize anions, stabilizing Q_B^- better than Q_B or QH. The free energy of protonating Q_B^- is 255 meV at pH 7 (Figure 3). This shifts the quinone pK_a' from 4.9 in solution to 2.7 in the protein (eq 9); therefore, the quinone will remain deprotonated as is found.

Table 2: MCCE-Calculated Redox Energies of Quinone in Proximal and Distal Q_B and Q_A Sites at pH 7 and E_h = 0 with P, the Bacteriochlorophyll Dimer, Neutral^a

half-reaction	$\Delta\Delta G_{\text{rxn}}$	ΔG_{pol}	(a) Q _B Proximal Site		$\Delta\Delta G_{\text{protein}}$	ΔG_{sol}	ΔG_{calc}	$\Delta\log K_d$
			ΔG_{nonel}	ΔG_{res}				
Q _B → Q _B ⁻	386	-389	0	-135	-138	145	7	-2.4
Q _B + e ⁻ + H ⁺ → Q _B H	-15	74	-23	-43	-7	269	262	-0.1
Q _B + 2e ⁻ → Q _B ⁻²	1471	-783	0	-533	155	340	495	2.6
Q _B + 2e ⁻ + H ⁺ → Q _B H ⁻	429	-342	-18	-179	-110	-1	-111	-1.8
Q _B + 2e ⁻ + 2H ⁺ → Q _B H ₂	18	117	-48	-38	49	-220	-171	0.8
Q _B ⁻ + e ⁻ → Q _B ⁻²	1085	-394	0	-398	293	195	488	5
half-reaction	$\Delta\Delta G_{\text{rxn}}$	ΔG_{pol}	(b) Q _B Distal Site		$\Delta\Delta G_{\text{protein}}$	ΔG_{sol}	ΔG_{calc}	$\Delta\log K_d$
			ΔG_{nonel}	ΔG_{res}				
Q _{BD} → Q _{BD} ⁻	317	-219	0	19	117	145	262	2.0
Q _{BD} + e ⁻ + H ⁺ → Q _{BD} H	-20	-16	-8	-15	-59	269	210	-1.0
Q _{BD} + 2e ⁻ → Q _{BD} ⁻²	1130	-433	0	-121	576	340	916	9.7
Q _{BD} + 2e ⁻ + H ⁺ → Q _{BD} H ⁻	365	-233	-8	-63	61	-1	60	1.0
Q _{BD} + 2e ⁻ + 2H ⁺ → Q _{BD} H ₂	15	-35	-18	8	-30	-220	-250	-0.5
half-reaction	$\Delta\Delta G_{\text{rxn}}$	ΔG_{pol}	(c) Q _A Site		$\Delta\Delta G_{\text{protein}}$	ΔG_{sol}	ΔG_{calc}	$\Delta\log K_d$
			ΔG_{nonel}	ΔG_{res}				
Q _A → Q _A ⁻	390	-254	0	-244	-108	145	37	-1.8
Q _A + e ⁻ + H ⁺ → Q _A H	-13	19	-15	-10	-19	269	250	-0.3
Q _A + 2e ⁻ → Q _A ⁻²	1479	-494	0	-588	397	340	737	6.7
Q _A + 2e ⁻ + H ⁺ → Q _A H ⁻	430	-219	-16	-8	-13	-1	-14	-0.2
Q _A + 2e ⁻ + 2H ⁺ → Q _A H ₂	18	13	-21	-12	-2	-220	-222	-0.03

^a ΔG_{sol} (E_h = 0, pH 7) is calculated as $2.3mk_bT(7 - \text{p}K_{\text{a,sol}}) - nF(0 - E_{\text{m,sol}})$ (eq 4) using $\text{p}K_{\text{a,sol}}$ and $E_{\text{m,sol}}$ from Table 1. For protonated quinones, $\Delta\Delta G_{\text{rxn}}$ and ΔG_{pol} are the values for the four conformers with different proton positions weighted by their occupancies in Monte Carlo sampling (Figure 5 and Table 4). $\Delta\Delta G_{\text{protein}}$ was calculated with eq 5, and ΔG_{res} was calculated with eq 10. (a) $\text{p}K_{\text{a}}'$ ($\text{p}K_{\text{a}}$ calculated from the energy of protonation at pH 7 (eq 9) for Q_B⁻, 2.7; for Q_B⁻², 17.3; and for Q_BH⁻, 8.0. (b) $\text{p}K_{\text{a}}'$ for Q_{BD}⁻, 7.9; for Q_{BD}⁻², 21.4; and for Q_{BD}H⁻, 12.2. (c) $\text{p}K_{\text{a}}'$ for Q_A⁻, 3.4; for Q_A⁻², 19.7; and for Q_AH⁻, 8.8.

Table 3: Energy of Protein Rearrangement on Quinone Reduction to Semiquinone in Three Binding Sites^a

half-reaction	ΔG_{res}	$\Delta G_{\text{res}}^{\text{prot}}$	$\Delta G_{\text{res}}^{\text{prot}*}$	$\Delta G_{\text{prot} \rightarrow \text{prot}*}^{\text{red}}$	$\Delta G_{\text{prot} \rightarrow \text{prot}*}^{\text{ox}}$
Q _A Q _B + e ⁻ → Q _A Q _B ⁻	-135	86	-318	-221	183
Q _A Q _{BD} + e ⁻ → Q _A Q _{BD} ⁻	19	69	-74	-50	93
Q _A Q _B + e ⁻ → Q _A ⁻ Q _B	-244	-263	-279	-19	35

^a ΔG_{res} was calculated with eq 10. Other terms were calculated with eq 11. $\Delta G_{\text{res}}^{\text{prot}}$ is the difference in interaction of the protein with quinone and semiquinone in the protein equilibrated around the quinone (prot), and $\Delta G_{\text{prot} \rightarrow \text{prot}*}^{\text{red}}$ is the energy it takes to move the protein into the conformation equilibrated around the semiquinone (prot*). The reduction could also happen by the alternate path, where first the protein is moved to the conformation equilibrated around the semiquinone requiring $\Delta G_{\text{prot} \rightarrow \text{prot}*}^{\text{ox}}$ and then the quinone is reduced in the pre-prepared protein, prot*, ($\Delta G_{\text{res}}^{\text{prot}*}$) (63).

Table 4: Position of Hydroxyl Proton on SerL223 and Proximal Q_B Found in Monte Carlo Sampling for the Different Q_B Redox States^a

redox state	Q _B				SerL223	
	H1	H2	H3	H4	A	B
Q _B					0.20	0.80
Q _B ⁻					0.00	1.00
Q _B ⁻²					0.00	1.00
Q _B H	0.54	0.16	0.00	0.30	0.56	0.44
Q _B H ⁻	0.08	0.72	0.00	0.20	0.07	0.93
Q _B H ₂	0.60	0.40	0.00	1.00	0.68	0.32

^a See Figure 5 for hydrogen locations. Monte Carlo sampling carried out at pH 7 with the quinone redox state fixed.

Second Reduction of Q_B. The two-electron quinone reduction is calculated to be 155 meV less favorable in the Q_B site than in solution (Table 2a). Q_B⁻² is destabilized 1471 meV by $\Delta\Delta G_{\text{rxn}}$, ≈ 4 times that found for Q_B⁻, while the favorable ΔG_{pol} is -783 meV, only twice that of the semiquinone. This is as expected given the linear response of the CE calculations used for the interaction of the static portions of the protein with the quinone charge. Thus, the pairwise interactions for groups with no conformation or ionization changes double as the charge on the quinone doubles. At the same time, the Born reaction field energy

increases as $\approx q^2$; therefore, it costs 4 times more energy to bury a charge of -2 than -1 (80). However, MCCE interactions with parts of the protein such as GluL212 and SerL223 that change do not show a linear response to changes in the charge. ΔG_{res} for Q_B + e⁻ → Q_B⁻ is much less favorable than $\Delta G_{\text{res}}^{\text{prot}*}$, because work is done to modify the Q_B site residues (eqs 10 and 11). However, there are only small changes in the position and ionization states of residues in the Q_B site on the second reduction. Only GluH173 becomes partially protonated to stabilize the dianion (Appendix I in the Supporting Information); therefore, little additional work is done to stabilize Q_B⁻². Thus, $\Delta G_{\text{res}}^{\text{prot}} \approx \Delta G_{\text{res}}^{\text{prot}*}$ for Q_B⁻ + e⁻ → Q_B⁻² and $\Delta G_{\text{res}}^{\text{prot}}$ for Q_B⁻ + e⁻ → Q_B⁻² is approximately twice $\Delta G_{\text{res}}^{\text{prot}*}$ for Q_B + e⁻ → Q_B⁻. This illustrates how the resultant interaction of the quinone with the site, ΔG_{res} , is largest when the site is pre-prepared ($\Delta G_{\text{prot} \rightarrow \text{prot}*}^{\text{red}} \approx 0$). However, even the added stabilization of the dianion because of the reaction occurring in a prepared protein is insufficient to pay for the much larger reaction field penalty. If the $E_{\text{m,sol}}$ is -195 mV, then the E_{m} for the second reduction of the anionic quinone (Q⁻ + e⁻ → Q⁻²) is -488 mV.

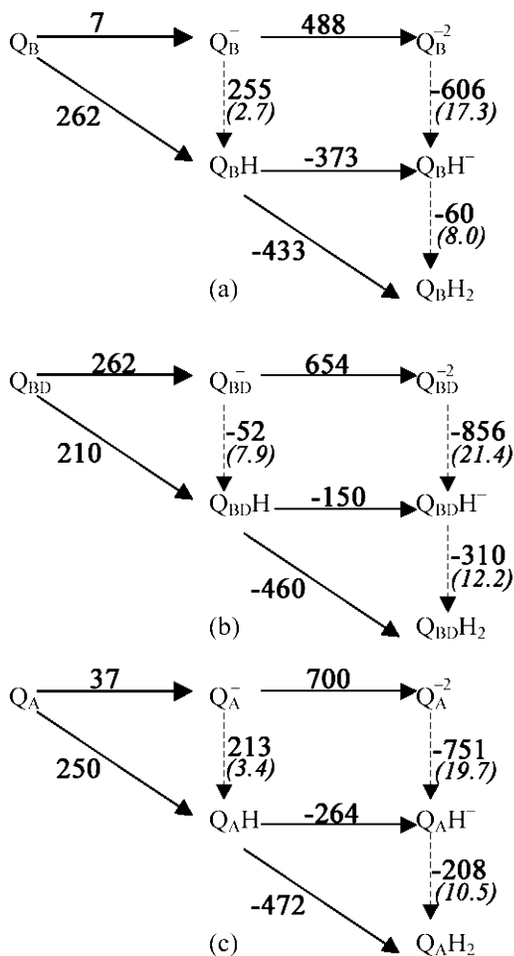


FIGURE 3: MCCE-calculated quinone redox energies in the three binding sites. (a) Q_B proximal site. (b) Q_{BD} distal site. (c) Q_A site. Horizontal arrows, electron transfers; dashed vertical arrows, proton transfers; diagonal arrows, proton-coupled electron transfers. ΔG values between all states connected by solid arrows were calculated by Monte Carlo sampling of the reactant and product states as a function of E_h . ΔG values of protonation reactions were obtained by closing the appropriated thermodynamic cycles. pK_a' values (in parentheses) were obtained from the ΔG of protonation at pH 7 using eq 9.

Protonation State of the Double-Reduced Q_B . The single-protonated fully reduced quinone, Q_BH^- , is stabilized by -111 meV, interacting with the protein in a manner similar to that found for the semiquinone (Table 2a). The ionization states in the acidic cluster are the same as with Q_B^- (Appendix I in the Supporting Information). The pK_a for binding the first proton to Q_B^{-2} is 12.7 in solution. The pK_a' is calculated to be pushed up to 17.3 in the Q_B site because of the destabilization of Q_B^{-2} (Table 2a and Figure 3).

The fully protonated, fully reduced Q_BH_2 is destabilized 49 meV by the protein. The interaction with the backbone (ΔG_{pol}) in particular with GlyL225 accounts for much of this unfavorable interaction. The pK_a of QH^- in solution is 10.7, but pK_a' is only 8.0 in the protein, a reflection of the destabilization of the second bound proton.

Location of Protons on Q_B . The quinone makes 2 hydrogen bonds in the Q_B site. One is to HisL190, which lies between the non-heme ferrous iron and Q_B . The other quinone carbonyl interacts with SerL223 and the backbone nitrogen of GlyL225. There are 4 positions for hydrogen sampled on Q_B (Figure 5). It can donate a hydrogen bond to (H1) or

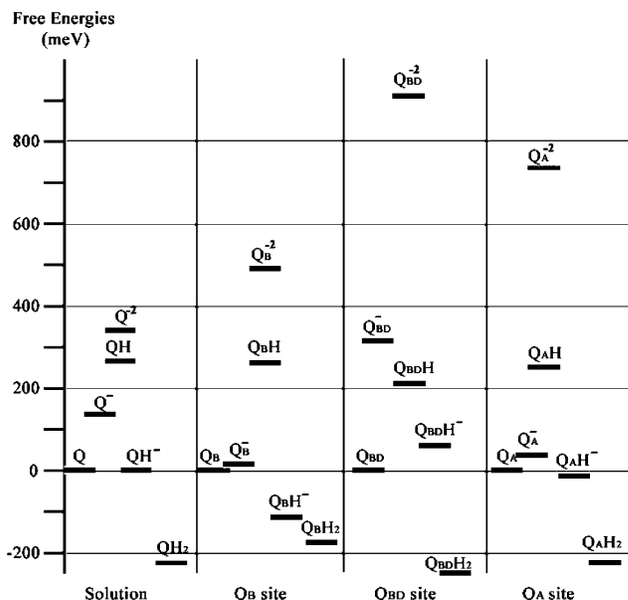


FIGURE 4: Energy levels of quinone products in Q_B , Q_{BD} , and Q_A sites. Values are from Table 2.



FIGURE 5: Available hydroxyl positions in different conformers of SerL223 and in QH and QH_2 states of ubiquinone. The UQ protons: H1, H2, H3, and H4. SerL223 protons: HA and HB. The side chain of the ligand HisL190 and the backbone of GlyL225 are also shown. The protein coordinates are taken from the 1AIJ structure (21) with the quinone moved into the 1AIG proximal position as described in the Materials and Methods.

accept a hydrogen bond from (H2) SerL223, or on the carbonyl proximal to the non-heme iron, it can point toward (H3) or away from (H4) HisL190. Two of the seven possible SerL223 conformers are occupied. In one, it is a donor to AspL213 and can be a hydrogen-bond acceptor from a protonated quinone (A), and in the other, it donates a hydrogen bond to Q_B (B). When Q_B is protonated, the position of protons on the Ser and the nearby quinone carbonyl are correlated to avoid clashes.

In the Q_B ground state, SerL223 donates a proton to Q_B (position B) with 80% probability (Table 4). This preference is strengthened in any of the anionic Q_B states. The SerL223 hydroxyl in the A position destabilizes Q_B^- by 67 meV, while the hydroxyl in the B position stabilizes it by -114 meV. A better test of the importance of the SerL223 position can be gained by comparing the E_m calculated with the Ser proton fixed, allowing the rest of the protein to come to equilibrium around each hydroxyl position. The E_m with SerL223 B is

−5 mV, essentially unchanged from that found in the free calculation. If the SerL223 is fixed in the A conformation, then the E_m is lowered to −95 mV. Thus, as proposed previously (81), if the SerL223 is oriented to donate a hydrogen bond to AspL213 and not to Q_B in the ground state, Q_B reduction by Q_A would be unfavorable.

In the neutral species Q_BH and Q_BH_2 , there is a distribution of quinone hydroxyl positions found in Monte Carlo sampling. It is somewhat more likely that the quinone will be a proton donor to SerL223. However, different positions are found because there are several competing energy terms. No individual hydrogen bond between neutral Q_B and SerL223 is stronger than 2 kcal/mol; therefore, multiple orientations have similar energies. The only disallowed quinone proton position is the one that clashes with the hydrogen from HisL190 (H3). Thus, Q_BH H2, where the quinone accepts a hydrogen bond from SerL223, has a better interaction with the backbone amide of GlyL225 and with SerL223 than H1, where the quinone donates a hydrogen bond to SerL223. However, H1 is found more often in Monte Carlo sampling because the correlated SerL223 position, A, has a more favorable interaction with AspL213. In addition, there is room for a proton in the region near HisL190 on the other carbonyl (H4), and this position is sufficiently favorable that it is partially occupied in Q_BH and Q_BH^- . Favorable Lennard–Jones interactions with HisL190 (ΔG_{nonel}) stabilize this position.

Relative Affinity of the Different Redox States of Quinone for the Q_B Site. The difference in interaction of the quinone and another, product redox state with the protein is provided by $\Delta\Delta G_{\text{protein}}$ for that reaction (Table 2). Thus, the binding energies of the two species differ by $\Delta\Delta G_{\text{protein}}$ (eq 12, Figure 2). For example, when Q_B and Q_B^- are compared, $\Delta\Delta G_{\text{protein}}$ is −138 meV, indicating that the semiquinone interacts 3.2 kcal/mol more favorably and will thus bind 220 times more tightly to the binding site (eq 12). This is in agreement with experimental data, showing that Q_B^- is more tightly bound than Q_B or Q_BH_2 (19). $\Delta\Delta G_{\text{protein}}$ is similar for Q_B^- or Q_BH^- and so their relative affinities for the binding site will be similar. Again, because $\Delta\Delta G_{\text{protein}}$ is similar for Q_B and Q_BH , there will be little difference in these K_d values (82). For Q_BH_2 , the $\Delta\Delta G_{\text{protein}}$ of 49 meV shows that the dihydroquinone is more stable in solution, keeping Q_BH_2 7 times less tightly bound than Q_B .

Redox Reactions in the Distal Q_B Site. There are different Q_B -binding sites found in X-ray crystal structures (12, 21). In wild-type RCs, two distinct Q_B -binding sites (distal and proximal to the non-heme iron) have been described. In the charge-separated $P^+Q_AQ_B^-$ structure (1AIG), the inner proximal semiquinone Q_B^- is located ≈ 5 Å closer to Q_A than it is in the dark-adapted PQ_AQ_B outer, distal site (1AIJ) and it has undergone a 180° propeller twist around the isoprene chain. In this outer site, the quinone is making hydrogen bonds with IleL224 and AlaL186.

The energetics of quinone reduction and protonation at the distal site (Q_{BD}) were calculated using the original distal, quinone position in 1AIJ. The E_m for the $Q_{BD} + e^- \rightarrow Q_{BD}^-$ half-reaction is −262 mV, while it is −7 mV in the proximal site (Table 2). The Q_B site stabilizes the semiquinone anion by −138 meV, while the Q_{BD} site destabilizes it by 117 meV (Table 2). When the two sites are compared, there is 80 meV less reaction field loss in Q_{BD} but 180 meV less favorable

ΔG_{pol} and 255 meV more unfavorable interactions with the rest of the protein. Reduction with residues in the protein equilibrated around the oxidized quinone, ΔG , is less unfavorable in the distal than the proximal site because the quinone is further from the acidic cluster of GluL212 and AspL213 (Table 3). However, the relaxation around the reduced quinone, $\Delta G_{\text{prot} \rightarrow \text{prot}^*}^{\text{ox}}$, provides far less stabilization. As in the proximal site, motions of SerL223 and protonation of GluL212 stabilize Q_{BD}^- . However, with an anionic semiquinone in the Q_{BD} site, GluL212 remains 70% ionized, destabilizing the anionic quinone by ≈ 90 meV (Table 2b, Appendix I in the Supporting Information).

The energies of the other redox states of Q_{BD} show that the distal site destabilizes anionic quinones, while the neutral protonated $Q_{BD}H$ and $Q_{BD}H_2$ are stabilized more than in the proximal Q_B site (Figures 3 and 4). There is adequate room in the binding site to protonate either carbonyl, although the position pointing toward the IleL224 HN is disallowed. Q_{BD}^{-2} is 421 meV higher in energy than Q_B^{-2} . The desolvation penalty for the fully reduced quinone is smaller in the distal site, but the stabilizing interactions with the backbone and other residues are much smaller. The dihydroquinone is however stabilized relative to the solution by −30 meV in the distal site, while it is destabilized by 49 meV in proximal site. Thus, QH_2 will be bound 3 times more tightly in the distal site than is the quinone (Table 2 and Figure 3).

Q_A Redox Reactions. In bacterial RCs, Q_A is tightly bound to the protein, and the Q_A position is well-defined in all crystal structures. The quinone is making hydrogen bonds to HisL219 and the backbone of AlaM260. The protein stabilizes reduction of Q_A by −108 meV relative to the quinone in solution (Table 2c). Given $E_{m,\text{sol}}$ for Q/Q^- of −145 mV, this results in an E_m of −37 mV close to the measured values of −45 (83) to −75 mV (75) at pH 7. $\Delta\Delta G_{\text{rxn}}$ destabilizes the anionic semiquinone by 390 meV, the same as found in the proximal Q_B site. Interactions with backbone dipoles stabilize reduction by −254 meV. The primary contributions are from amides M260–271, in the loop leading into and the first 10 residues of the E transmembrane helix. ΔG_{res} is −244 meV. Reduction of Q_A is favored even in the protein equilibrated with the oxidized quinone ($\Delta G_{\text{res}}^{\text{prot}}$) (Table 3), in agreement with the ability of the dark adapted, frozen protein to form Q_A^- following activation of the RCs with light (84). $\Delta G_{\text{prot} \rightarrow \text{prot}^*}^{\text{ox}}$ is 35 meV, showing there are only small rearrangements in the MCCE calculation on Q_A reduction. Several polar residues including ThrM261, ThrM222 and TyrH40 rearrange their hydroxyl dipoles to stabilize the charge. The resultant $\Delta\Delta G_{\text{protein}}$ of −108 meV indicates that Q_A^- will be bound 72 times tighter than Q_A .

There are a variety of studies that suggest a linkage between Q_A and Q_B (1, 85). The E_m for Q_A is slightly less negative when Q_B is reduced than when it is oxidized (Table 5a). The charge–charge interaction between Q_A^- and Q_B^- itself lowers the E_m by ≈ 55 mV. However, neutralization of GluL212 in the Q_B -site acidic cluster raises the potential by ≈ 62 mV. The resultant E_m thus changed by only ≈ 10 mV when Q_B is reduced.

As in the proximal Q_B site, the Q_A site stabilizes species with a −1 charge relative to the neutral or doubly reduced

Table 5: Selected Quinone Half-Reactions Calculated at pH 7

half-reactions ^a	$E_{m,sol}$ (mV)	ΔG_{sol} (meV)	$\Delta G_{protein}$ (meV)	E_m (mV)	ΔG (meV)
Q_AQ_B + e⁻ → Q_A⁻Q_B	-145	145	-108	-37	37
<i>Q_AQ_B⁻ + e⁻ → Q_A⁻Q_B⁻</i>	-145	145	-121	-24	24
Q_AQ_BH + e⁻ → Q_A⁻Q_BH	-145	145	-113	-32	32
<i>Q_AQ_B + e⁻ → Q_AQ_B⁻</i>	-145	145	-137	-7	7
<i>Q_A⁻Q_B + e⁻ → Q_A⁻Q_B⁻</i>	-145	145	-144	-1	1
<i>Q_AQ_B⁻ + e⁻ → Q_AQ_B⁻²</i>	-195	195	295	-490	490
Q_AQ_BH + e⁻ → Q_AQ_BH⁻	270	-270	-103	373	373
Q_AQ_B + e⁻ + H⁺ → Q_AQ_BH		269	-7		262
Q_A⁻Q_B + e⁻ + H⁺ → Q_A⁻Q_BH		269	-10		259
<i>Q_AQ_B⁻ + e⁻ + H⁺ → Q_AQ_BH⁻</i>		-146	24		-122
<i>Q_AQ_BH + e⁻ + H⁺ → Q_AQ_BH₂</i>		-488	56		-433

reactions ^b	derived from	ΔG (meV)	exp. ΔG (meV)
Q_AQ_B + e⁻ → Q_A⁻Q_B	ΔG_1^a	37	45 (83) to 75 (75)
<i>Q_A⁻Q_B → Q_AQ_B⁻</i>	$\Delta G_4 - \Delta G_1$	-30	-70 (86-88)
<i>Q_AQ_B⁻ + e⁻ → Q_A⁻Q_B⁻</i>	ΔG_2^c	24	20 (75)
<i>Q_A⁻Q_B⁻ → Q_AQ_B⁻²</i>	$\Delta G_6 - \Delta G_2$	464	>240 ^d
<i>Q_A⁻Q_B⁻ + H⁺ → Q_A⁻Q_BH</i>	$\Delta G_9 - \Delta G_5$	258	150 (90) ^e
<i>Q_AQ_B⁻² + H⁺ → Q_AQ_BH⁻</i>	$\Delta G_{10} - \Delta G_6$	-620	>-215 ^d
<i>Q_A⁻Q_BH → Q_AQ_BH⁻</i>	$\Delta G_7 - \Delta G_3$	-405	-250 ± 40 (90)
<i>Q_AQ_BH⁻ + H⁺ → Q_AQ_BH₂</i>	$\Delta G_{11} - \Delta G_7$	-60	-90 ± 18 (89, 92) ^f

^a MCCE-calculated redox midpoint potential of selected half-reactions involving reduction of the proximal Q_B. The quinone that is undergoing the redox reaction is in bold. An unchanged, charged quinone is italicized. ^b Calculated free energy of the electron- and proton-transfer reactions in bacterial RC. ΔG values derived from appropriate half-reactions (Table 5a and Figure 6). ^c ΔG calculated for the reaction at $E_h = 0$; therefore, $\Delta G = -nFE_m$. ^d Personal communication with C. A. Wraight. ^e ΔG at pH 7 derived from the estimated ΔG of 180 ± 30 meV at pH 7.5 (90). ^f ΔG at pH 7 derived from the measured pK_a of 8.5 ± 0.3 (89, 92).

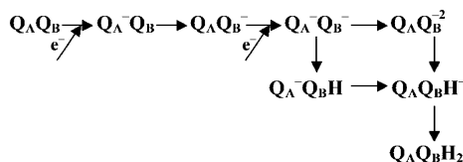


FIGURE 6: Electron- and proton-transfer reactions in bacterial RCs. The free energy of the proton transfer $Q_A^-Q_B^- + H^+ \rightarrow Q_A^-Q_BH$ is calculated from the free-energy difference between the reactions $Q_A^-Q_B^- + H^+ \rightarrow Q_A^-Q_BH$ and $Q_A^-Q_BH \rightarrow Q_AQBH^-$.

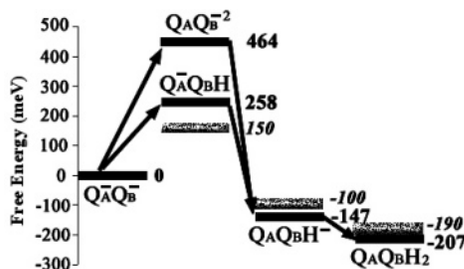


FIGURE 7: Comparison of calculated and experimental energy levels for doubly reduced states. Solid lines, the MCCE-calculated free energies (Table 5b); and gray lines, experimental energy values (see Table 5b for details).

forms of the quinone. Protonation of Q_A is more favorable than Q_B with a pK_a' of 3.4. There is adequate room in the binding site to protonate either carbonyl, although protonation of the carbonyl near AlaM260 is favored. The site destabilizes the dianion by 397 meV (Table 2c), significantly more than the Q_B site.

Energetics of the Electron Transfers between Q_A and Q_B:
First Electron Transfer, Q_A⁻Q_B → Q_AQ_B⁻. In RCs, Q_A is reduced forming Q_A⁻Q_B, and then the electron is transferred to Q_B yielding Q_AQ_B⁻ (Figure 6). The MCCE calculated E_m values for Q_AQ_B/Q_A⁻Q_B and Q_AQ_B/Q_AQ_B⁻ half-reactions are -37 and -7 mV (Table 5a). The free-energy difference, ΔG , for Q_A⁻Q_B → Q_AQ_B⁻ is -30 meV (Table 5b), close to the

experimental results of ≈ -70 meV determined from the rate of P⁺Q_B⁻ charge recombination, which proceeds via an equilibrated, thermal reduction of Q_A (86-88). Any uncertainties in $E_{m,sol}$ for Q/Q⁻ cancel in the calculated ΔG . Q_A and Q_B are arranged with c₂ symmetry around the non-heme iron. The two sites have similar $\Delta\Delta G_{rxn}$. However, Q_B⁻ has a much more favorable interaction with the backbone amide dipoles, ΔG_{pol} , while Q_A⁻ has a more favorable interaction with the residues of the protein (Table 2). The result is that, following the rearrangements of the Q_B site, the first electron transfer from Q_A⁻ to Q_B is favorable.

Second Electron Transfer and Subsequent Protonation, Q_A⁻Q_B⁻ → Q_AQ_BH₂. The second electron transfer results in the state Q_AQ_BH⁻ (Figure 6) (29, 89). A mechanism where Q_B⁻ is protonated forming Q_BH, followed by the second electron transfer, is supported by the free energy and pH dependence of the rate of the reduction of Q_B⁻ by Q_A⁻ (16). This conclusion is also supported by the analysis of the pH dependence of rhodoquinone (RQ) reduction in the Q_B site. The pK_a of RQ in solution is 7.3 ± 0.2, binding a proton more tightly than UQ (90). The pH dependence of the rate of the second electron transfer shows that RQ_BH is an intermediate when RQ is substituted for UQ. The free energy calculated here for Q_A⁻Q_B⁻ → Q_AQ_B⁻² of 464 meV versus Q_A⁻Q_B⁻ → Q_A⁻Q_BH of 258 meV supports the mechanism with proton transfer before electron transfer at pH 7 (Figure 7). These results are also consistent with prior electrostatic calculations in *Rps. viridis* RCs (91).

The free-energy change for Q_A⁻Q_B⁻ → Q_A⁻Q_BH was estimated to be 180 meV at pH 7.5 from a Marcus analysis comparing the electron-transfer rates when RQ and UQ are in the Q_B site (90). This would give a ΔG of 150 meV at pH 7 and a pK_a' for Q_B⁻ of 4.5. It is calculated to require 258 meV to protonate Q_B⁻, 110 meV more than the value derived from the experiment. The calculated pK_a' for Q_B⁻ → Q_BH is 2.7. The unfavorable energy for the protonated

semiquinone could arise from the proton position not being properly optimized in the calculations or from the negative charge on the semiquinone being overly stabilized. However, a high energy for Q_BH is not ruled out by the observed rate for the second reduction of Q_B . The rate is proportional to the product of the concentration of $Q_A^-Q_BH$ and the rate of electron transfer from Q_A^- to Q_BH (12). If the $Q_A^-Q_BH$ level is 110 meV higher as suggested here, its concentration will be 70 times smaller. To maintain the same $Q_A^-Q_B^- \rightarrow Q_A^-Q_BH$ reaction rate [$k_{AB}^{(2)} \approx 10^3 \text{ s}^{-1}$ (92)], the electron-transfer rate would have to be 70 times faster than the previous estimate of $\approx 10^6 \text{ s}^{-1}$ (12, 90).

Kinetic measurements provide a free energy for the reaction $Q_A^-Q_B^- + e^- + H^+ \rightarrow Q_AQ_BH^-$ of $-70 \pm 10 \text{ meV}$ at pH 7.5 (89), equivalent to a ΔG of -100 meV at pH 7. ΔG for $Q_A^-Q_BH \rightarrow Q_AQ_BH^-$ would then be -250 meV . The value calculated here is -405 meV , significantly more favorable. One possible problem is that the E_m for Q_BH^- relies on the $E_{m,\text{sol}}$ for Q^-/Q^{2-} . A higher energy of Q^{2-} would raise that of $Q_AQ_BH^-$, reducing the calculated ΔG between $Q_A^-Q_B^-$ and $Q_AQ_BH^-$.

Given the high energy of the $Q_AQ_B^{2-}$ state, the free-energy difference between $Q_A^-Q_B^-$ and $Q_AQ_B^{2-}$ states is not measurable. The E_m (Q_B^-/Q_B^{2-}) has been estimated to be more negative than -240 mV (C. A. Wraight, personal communication). The calculated value is -464 mV . An experimental pK_a of >10.7 was obtained for binding a proton to Q_B^{2-} in the state $Q_A^-Q_B^{2-}$ (36, 93). The calculated value is 17.3. The pK_a for $Q_AQ_BH^- + H^+ \rightarrow Q_AQ_BH_2$ was estimated to be 8.5 ± 0.3 from the steady-state proton uptake by Q_B^{2-} (89, 92). This corresponds to a reaction free energy of $-90 \pm 18 \text{ meV}$ at pH 7, in reasonable agreement with the calculated value of -60 meV .

DISCUSSION

Quinone Energies in the Three RC-Binding Sites. In RCs of *Rb. sphaeroides* and *Rps. viridis*, three quinone-binding sites are found in various X-ray structures (21, 82, 94, 95). The two quinones symmetrically arranged around the non-heme iron are clearly identified with functionally well-characterized Q_A and Q_B (1, 2, 12). Q_A is the first quinone reduced, accepting an electron from the bacteriopheophytin in $\approx 150 \text{ ns}$ (96). On two turnovers, Q_A^- reduces Q_B and then the Q_B semiquinone with rates of $\approx 100 \mu\text{s}$ (14, 97) prior to Q_B leaving the protein as dihydroquinone (17). The role of the distal quinone site is not yet clear (21–24, 26). MCCE has been used here to see how the RC structure influences the free energy of the quinone redox states so that ubiquinone can carry out its different functions in these sites (Figure 4 and Table 2). Different states are stabilized so that in each site the appropriate reactions are energetically accessible. In addition, each quinone species has a different affinity for each binding site. For example, quinone comes and goes in Q_B and Q_{BD} sites but never leaves the Q_A site. The connection between the stabilization of a redox state and the relative affinity of different redox states shows why the semiquinone does not bind to the Q_{BD} site but does not leave the Q_B site. (Figure 2) (98).

Q_A . Q_A is reduced by the bacteriopheophytin anion, which only lives for a few nanoseconds. Thus, this quinone never dissociates so that it is always ready to react (74, 99). Q_A

serves as a single-electron acceptor cycling between Q_A and Q_A^- without binding protons. The MCCE calculations show that the Q_A site stabilizes only the semiquinone (Table 2). The calculated E_m , made assuming an $E_{m,\text{sol}}$ for ubiquinone in water of -145 mV , is -37 mV , in good agreement with the measured value of $\approx -60 \text{ mV}$ (75, 83). The protein interacts weakly with a proton on either carbonyl. The free energy of proton binding is 89 meV less favorable in the Q_A site than in solution, lowering the semiquinone pK_a' from 4.9 to 3.4.

Q_B . Q_B has been measured to be 20 times more weakly bound than Q_A (74, 99). This is acceptable because Q_A^- lives for hundreds of milliseconds; therefore, Q_B has time to bind to an empty site before $P^+Q_A^-$ returns to the ground state, which would waste the energy of the absorbed photon. Q_B cycles through the Q_B , Q_B^- , Q_BH , Q_BH^- , and Q_BH_2 states during turnover, alternating reduction and protonation reactions. The proximal Q_B site stabilizes the anionic states Q_B^- and Q_BH^- and interacts weakly or slightly unfavorably with the protonated neutral states Q_BH and Q_BH_2 . An important requirement for RC function is that Q_A^- can reduce both quinone and semiquinone species within the millisecond lifetime of Q_A^- . Despite the c_2 symmetry of Q_A and Q_B sites in the protein, the Q_B site stabilizes the semiquinone somewhat more than the Q_A site, in large part because of the larger positive potential from the backbone dipoles in the proximal Q_B site. The free energy for the first electron transfer is modest, $\approx -70 \text{ meV}$ experimentally (86–88) and -30 meV here; therefore, Q_B^- is not deeply trapped. This allows the electron to return to P^+ reforming the ground state via reduction of Q_A with a half-time of $\approx 1 \text{ s}$ if a second electron is not delivered to Q_B . The Q_B site stabilizes Q_BH a little relative to Q in solution (Figure 4), making the ΔG between Q_B^- and Q_BH larger than it is in solution. However, the site destabilizes Q_B^{2-} more; therefore, Q_BH and not the dianion is the intermediate for the second reduction of Q_B , in agreement with previous suggestions (90, 100). If the semiquinone species were to leave the Q_B site before the second electron is delivered from Q_A^- , the energy of the photon would be lost. The -140 meV stabilization of Q_B^- and -110 meV stabilization of Q_BH^- ensures that these will bind more tightly to the Q_B site than does the quinone. The Q_B site stabilizes anions without a strong penalty for proton binding. This allows a favorable free energy for electron transfer from Q_A^- to Q_BH to form Q_BH^- . There is a modest, 50 meV , destabilization of Q_BH_2 . This puts the energy of Q_BH_2 below that of Q_BH^- but above that of QH_2 in solution, yielding favorable proton binding to Q_BH^- but weak binding to the Q_B pocket.

Q_{BD} . RC crystal structures have shown that ubiquinone binds in several positions in the Q_B site (95). X-ray crystal structures were compared of RCs frozen in the light and dark. In the dark-adapted structures, where the quinone was assumed to be oxidized, it was found in the outer distal site (21). In the light-adapted structure, where the state was assumed to be $P^+Q_B^-$, the quinone is in the proximal site. The distal headgroup moves $\approx 5 \text{ \AA}$ out of the protein into a location overlapping the tail of a proximal quinone. In addition, the two positions differ by a 180° rotation around the isoprene tail. Prior kinetic data had shown that the electron transfer from Q_A^- to Q_B is gated by a conformational change (101). The motion from proximal to distal binding

sites was proposed as a good candidate for this rate-determining step (21). However, several more recent experiments have cast doubt on this hypothesis. Recent time-resolved crystallography in *Rps. viridis* (26) shows that only the proximal site is occupied at room temperature in ground and $P^+Q_B^-$ states of RCs that are undergoing turnover. This agrees with FTIR studies that there are no changes in the quinone position in active *Rb. sphaeroides* RCs (22, 23). Likewise, the kinetic studies show that the tail has little effect on the electron transfer from Q_A^- to Q_B , unexpected if the quinone needs to undergo rotation and translation in the rate-determining step (15, 24, 99). The quinone position in crystals is sensitive to mutation (102), temperature, and cryoprotectant (103). Computational analysis has explored the transition from the proximal to distal position showing there is a little barrier to movement without (40, 41) and a large barrier to movement with (39) the 180° rotation. The protonation of the acidic cluster changes the relative affinity of the proximal and distal sites (40, 41). In addition, as is found here (Appendix I in the Supporting Information), the proton uptake on Q_B reduction depends on the quinone location (27).

MCCE analysis shows that the distal Q_B site (Q_{BD}) destabilizes all anionic quinone species while stabilizing the protonated neutral states (Figure 3). Both ΔG_{pol} and ΔG_{res} are significantly less favorable than they are in the Q_B site. Reduction of Q_{BD} to Q_{BD}^- is ≈ 120 meV less favorable than it is in solution and 260 meV less favorable than in the proximal Q_B site. Thus, Q_A could not reduce Q_{BD} ; therefore, it is highly unlikely that electron transfer could occur to a quinone bound there. This is in agreement with prior calculations of RCs *Rps. viridis* (44). Earlier calculations on *Rb. sphaeroides* RCs show a 210 meV difference between reduction at the two sites, in good agreement with the values found here (38, 46). However, the neutral, protonated quinone species $Q_{BD}H$ and $Q_{BD}H_2$ are stabilized in the distal binding site. Thus, the dihydroquinone will be bound ≈ 3 times more tightly than the quinone to the distal site. It has been suggested that dihydroquinone, reduced during data collection, is the species found in the RC crystal Q_{BD} site (55, 94).

Comparisons with Earlier Experimental and Computational Studies. The values have been presented to the nearest millivolt here as an aid for tracking the connections between numbers in different tables. However, neither the calculations nor the experimental data to which it is compared are known with this certainty. The uncertainty of the Monte Carlo sampling for this model of the RCs is less than 5 mV. MCCE analysis of benchmark E_m (63) and pK_a (54) values show that >90% of the residues have an error of less than 60 mV or 1 pH unit. The calculations are sensitive to the starting crystal structures, parameters, and conformers (53). MCCE does not allow the backbone to change conformation, and only limited side-chain positions are sampled.

Ionization of the Acidic Cluster. Perhaps the most difficult part of the simulation to define is the protonation state of the acidic cluster near Q_B , including GluL212, AspL213, AspL210, and GluH173 (Appendix I in the Supporting Information). Their ionization states when the quinone is in the ground and Q_B^- states have been the subject of many experimental (1, 2, 33) and computational (37, 38, 45, 46) studies. Experiment, especially FTIR measurements, have

indicated that a partially deprotonated GluL212 becomes protonated when Q_B is reduced (35, 104, 105). Simulations find that GluL212 and AspL213 are strongly coupled. Some calculations find that GluL212 is ionized (38, 43, 45, 46), and others find that AspL213 is ionized (37, 41). In calculations, the pair usually have a net charge of -1 as they do here. When Q_B is reduced, GluL212 and AspL213 are always neutral, retaining a cluster net charge of -1 .

In the simulations starting with 1AIJ presented here, when Q_B is oxidized, GluL212 is ionized (Appendix I in the Supporting Information). In parallel, in MCCE calculations carried out with 1M3X (106), GluL212 is neutral and AspL213 is ionized in the ground state (data not shown). This ease of shifting of the order of acid ionization arises because these two acids interact strongly; therefore, only one can be ionized. They have similar pK_a values in the absence of the interaction with each other; therefore, their ionization free energy is similar. Modest changes in the structure cause one or the other to be ionized first, keeping the other neutral to high pH. However, the E_m for Q_B differs by only 10 mV, and the proton uptake is the same ($0.7 H^+/e^-$) in the simulations on 1AIJ and 1M3X. This is a result of the free energy of the reactant microstates, with $Q_B/L212^-/L213H$ and $Q_B/L212H/L213^-$ being very close together. The product state is always $Q_B^-/L212H/L213H$. This explains why the free energy of electron transfer from Q_A^- to Q_B in our earlier calculations, where GluL212 was protonated and AspL213 was ionized in the ground state, differ by only ≈ 40 meV from those presented here (37). Similar clusters of acidic residues with coupled protonation are also found in bacteriorhodopsin (65).

Importance of the Membrane. While most measurements on bacterial RCs are carried out with detergent-solubilized protein, RCs function *in vivo* embedded in the cell membrane. The reported reaction E_m and pK_a values are somewhat dependent on the measurement conditions. For example, quinone E_m values are lower in chromatophores than in the isolated RC (107). In contrast, preliminary calculations show that addition of a low dielectric slab to simulate the membrane (108) raises both the Q_A and Q_B semiquinone E_m values because there are more basic residues further from the quinones and more acidic residues nearby. Without the low dielectric slab, the surrounding water screens the influence of the more distant positively charged groups. Experiments show addition of a few bound cardiolipin molecules can lower the $Q_A/Q_A^- E_m$ by -30 mV; therefore, at least some of the difference found in membranes may be due to the binding of specific lipids rather than the impact of dielectric screening by the membrane (109). Measurements provide a ΔG for electron transfer from Q_A^- to Q_B of ≈ 65 and ≈ 75 meV in chromatophores when P is neutral (110). In chromatophores, the reaction is 45 meV more favorable when P is oxidized, indicating P^+ stabilizes Q_B^- more than Q_A^- (110). In contrast, the ΔG for electron transfer from Q_A^- to Q_B is relatively independent of the ionization state of P in isolated RCs (87, 111). The quinone pK_a values can also depend on the environment. Thus, an experimental estimate for the pK_a of $Q^- + H^+ \rightarrow QH$ is 4.5 in detergent-solubilized RCs (90) and 6 in chromatophores (107).

Role of SerL223. The position of the hydroxyl on SerL223 depends on the initial proton distribution on the acidic cluster in the Q_B site. When AspL213 is neutral, as in these 1AIJ

calculations, two hydroxyl positions are found in Monte Carlo sampling; therefore, in the ensemble, some SerL223 donate a hydrogen bond to and others accept a hydrogen bond from the neutral Q_B (Table 4). However, if AspL213 is ionized when Q_B is neutral, as it is in the 1M3X calculations, the hydroxyl of SerL223 points to the Asp, away from Q_B . In both cases, when Q_B is reduced, SerL223 serves as a hydrogen-bond donor to the anionic semiquinone. The results presented here are in good agreement with a computational analysis that focused on the importance of SerL223 (81). The results presented here reaffirm the dependence of the Ser hydroxyl orientation on the charge of AspL213 and the approximate magnitude of the change in $Q_B E_m$ if the Ser is fixed in one position or the other. If AspL213 is neutral when Q_B is oxidized, the conformations with different proton positions are close in energy and small changes in structure or fitting parameters could easily change the distribution. Thus, it seems likely that the Ser hydroxyl reorientation provides a significant barrier to electron transfer from Q_A^- to Q_B only if AspL213 and not GluL212 is the member of the acidic cluster that is ionized when Q_B is oxidized.

Pathway of Proton Transfer to Q_B . Site-directed mutagenesis suggests that the first proton is transferred to the carbonyl from AspL213 via SerL223 and that the second proton is transferred to oxygen near HisL190 through a pathway involving GluL212 (29, 112, 113). This is consistent with the first quinone proton being bound to the carbonyl near the SerL223 (Table 4). However, in these calculations, the proton on the carbonyl hydrogen-bonded to HisL190, pointing away from the His NE2 hydrogen, and the non-heme iron is found in 30% of Q_BH (Figure 5). This position has weak but favorable Lennard–Jones interactions with both HisL190 and the non-heme iron. It may be that this proton can compete with protonation of the other carbonyl in equilibrium simulations but would not be seen in RCs because protonation from the Ser is much faster. It also maybe that the favorable nonelectrostatic interactions are over-estimated.

Importance of $E_{m,sol}$. The calculated E_m values versus S.H.E., quinone pK_a values, and reaction ΔG values (Table 5) depend on the reference $E_{m,sol}$ and $pK_{a,sol}$ (63). As described in the Materials and Methods, aqueous E_m and pK_a values for ubiquinone have an uncomfortable number of uncertainties. Different errors in $E_{m,sol}$ for different half-reactions will yield errors in calculated ΔG values. For example, the calculated free-energy change for $Q_A^-Q_BH \rightarrow Q_AQ_BH^-$ is 150 meV more favorable than that estimated from the measurements (Figure 7) (89). The calculated ΔG for the reaction relies on $pK_{a,sol}$ for Q^{-2} and $E_{m,sol}$ for the reduction of the semiquinone, as well as the calculated shift of the energy in the protein, $\Delta\Delta G_{protein}$. Errors in any of these terms will change the results.

Different analyses of RC electrochemistry have used different values for $E_{m,sol}$. For example, Knapp and co-workers (38) use -360 mV for the $E_{m,sol}$ for Q/Q^- in DMF, a noninteracting reference solvent. In contrast, we prefer that all solution values are estimated in aqueous solution, because continuum calculations are optimized to determine transfers from water to the protein. Thus, the aqueous $E_{m,sol}$ of -145 mV, used here, is 215 mV more positive than that found in DMF. The E_m values from ref 38 are reported to be more

negative than those obtained here. However, their shifts in E_m because of the protein actually favor Q_A or Q_B reduction to semiquinone by ≈ 100 meV more than those found here. The ΔG for the electron transfer between the two quinones is a value that is independent of $E_{m,sol}$. Calculated values at pH 7 range from -65 meV (37) to -60 meV (38) to -30 meV here. This is a remarkable agreement considering the complexity of these calculations.

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SUPPORTING INFORMATION AVAILABLE

Appendix I, protonation and ionization states of selected residues at pH 7 calculated by MCCE calculation here and from previous calculations and experiments. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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