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Biochimica et Biophysica Acta



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Review

The Q-cycle reviewed: How well does a monomeric mechanism of the bc_1 complex account for the function of a dimeric complex?

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ARTICLE INFO

Article history: Received 6 February 2008 Received in revised form 26 March 2008 Accepted 23 April 2008 Available online 1 May 2008

Keywords Q-cycle Constraints on molecular mechanism bc_1 complex Kinetic model Coulombic interaction Thermodynamic model

ABSTRACT

Recent progress in understanding the Q-cycle mechanism of the bc_1 complex is reviewed. The data strongly support a mechanism in which the O₀-site operates through a reaction in which the first electron transfer from ubiquinol to the oxidized iron-sulfur protein is the rate-determining step for the overall process. The reaction involves a proton-coupled electron transfer down a hydrogen bond between the ubiquinol and a histidine ligand of the [2Fe-2S] cluster, in which the unfavorable protonic configuration contributes a substantial part of the activation barrier. The reaction is endergonic, and the products are an unstable ubisemiquinone at the Q₀-site, and the reduced iron-sulfur protein, the extrinsic mobile domain of which is now free to dissociate and move away from the site to deliver an electron to cyt c_1 and liberate the H⁺. When oxidation of the semiquinone is prevented, it participates in bypass reactions, including superoxide generation if O_2 is available. When the *b*-heme chain is available as an acceptor, the semiquinone is oxidized in a process in which the proton is passed to the glutamate of the conserved -PEWY- sequence, and the semiquinone anion passes its electron to heme $b_{\rm L}$ to form the product ubiquinone. The rate is rapid compared to the limiting reaction, and would require movement of the semiquinone closer to heme $b_{\rm L}$ to enhance the rate constant. The acceptor reactions at the Qi-site are still controversial, but likely involve a "two-electron gate" in which a stable semiguinone stores an electron. Possible mechanisms to explain the cyt b_{150} phenomenon are discussed, and the information from pulsed-EPR studies about the structure of the intermediate state is reviewed.

The mechanism discussed is applicable to a monomeric bc_1 complex. We discuss evidence in the literature that has been interpreted as shown that the dimeric structure participates in a more complicated mechanism involving electron transfer across the dimer interface. We show from myxothiazol titrations and mutational analysis of Tyr-199, which is at the interface between monomers, that no such inter-monomer electron transfer is detected at the level of the b_L hemes. We show from analysis of strains with mutations at Asn-221 that there are coulombic interactions between the *b*-hemes in a monomer. The data can also be interpreted as showing similar coulombic interaction across the dimer interface, and we discuss mechanistic implications.

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1. Introduction

The ubiquinol:cytochrome c oxidoreductase (bc_1 complex) family of redox-linked proton pumps form the core of all major electron transfer chains, with an ancestry dating back to before the great divide. Their importance needs no further emphasis beyond their role in carrying the energy flux of the biosphere. It is generally agreed that

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a Q-cycle mechanism accounts for the activity of the enzyme, but there are many variants, and which particular variant best explains the finer points is still under debate [1–11]. In the simpler bacterial systems, the kinetic and thermodynamic behavior is well described by the modified Q-cycle [12,13]. This has been discussed extensively in recent reviews [2–5,11,13], and is shown schematically in Fig. 1. The *bc*₁ complex has a dimeric structure incorporating 4 metal centers in each monomer (hemes *b*_H and *b*_L in cytochrome (cyt) *b*, heme *c*₁ in cyt *c*₁, and a [2Fe–2S] cluster in the iron–sulfur protein (ISP)), which provide electron transfer chains connecting two quinone processing sites, and three additional catalytic interfaces. Despite this complexity, the mechanism is better understood than that for other respiratory complexes. The oxidation of ubiquinol (QH₂) occurs at the Q_o-site (also

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Fig. 1. The modified Q-cycle mechanism. The mechanism demonstrated in the 1980's, superimposed on the *Rb. sphaeroides* structure (PDB file 2QJK). Electron transfers are shown by blue arrows; H⁺ release or uptake by red arrows, exchange of Q or QH_2 by open blue-green (Q_o -site) or yellow (Q_i -site) arrows; inhibition sites by cyan arrows. See text for a description of the operation.

called the Q_P-site), where the two electrons are diverted down two different chains. The first electron goes down the high-potential chain of ISP, heme c_1 , and cyt c (or c_2 in bacteria), to an oxidant (a cytochrome oxidase in respiratory chains, or the oxidized reaction center in photosynthetic systems). Removal of one electron leaves a semiquinone (SQ) at the Q₀-site. The second electron is delivered to a low-potential chain of hemes $b_{\rm L}$ and $b_{\rm H}$, which passes the electron on to the Q_i -site (Q_N -site), where it reduces either ubiquinone (Q) to SQ, or SQ to QH₂. Because reduction of Q at the Q_i-site requires two electrons overall, the bifurcated reaction at the Q₀-site has to occur twice to provide these, and to complete a full turnover of the enzyme. The location of the two Q-sites on opposite sides of the membrane gives rise to an electrogenic flux through the low-potential chain, which provides the electrogenic arm, while diffusion of Q and QH₂ across the membrane with oxidation of 2QH₂ (with release of 4H⁺), or reduction of Q (with uptake of 2H⁺) provides the neutral arms of a Mitchellian proton-pumping loop.

The ancestral history of the bc_1 complex has left an unfortunate mechanistic legacy [14–18]. Oxidation of ubihydroquinone (ubiquinol, QH₂) at the Q_o-site of the complex operates through an intermediate semiquinone (SQ) that has a high reactivity with O₂. This leads to production of superoxide anion, the precursor of reactive oxygen species (ROS) that play havoc with the cellular machinery, and ultimately lead to cellular suffocation through damage to mitochondrial DNA [18,19]. There is some controversy over what fractional contribution this reaction makes to the overall cellular ROS production in humans, but a substantial literature suggests that it may be the major source under some physiological conditions and in some genetic pathologies [20–23]. This design-defect likely reflects an "invention" when the biosphere was anaerobic, and one of the interests in studying the complex has been to see how evolution has subsequently honed the architecture to minimize the effect.

Much of the critical work has investigated the bc_1 complex in *Rhodobacter* species of photosynthetic bacteria. The major advantage of working in photosynthetic systems lies in the fact that the bc_1 complex is tightly integrated into the photosynthetic machinery [24], so that excitation with a saturating flash generates the substrates stoichiometrically, allowing exploration of the kinetics of a single turnover of the intact apparatus on the microsecond time scale. From this work, a detailed description of the mechanism has been proposed, in which the partial reactions have been dissected out, and characterized through kinetic and thermodynamic parameters sufficient to account for the bulk of available experimental data [4,5,25,26]. With the availability of structures [27-33], this physicochemical description has been fleshed out in molecular detail. Although the structures provided strong support for the modified Q-cycle, they also introduced some novel features [34-36]. The structures have also put much detailed work using mutagenesis on a firmer basis, providing insights that have allowed an atomic-level analysis of mechanism.

In this paper, we briefly review recent developments that have provided us with an economical model for turnover in a monomeric complex, and then explore the question whether this is sufficient to explain turnover of the dimer.

1.1. Mobility of the ISP ectodomain

Recognition of the mobility of the extrinsic domain [28] launched a period of controversy about the role of movement of the ISP in control

of function. Much of the speculation has focused on a possible controlling role of such movement. From our own work [36,37], and from Millet's results [38,39] it is clear that movement is always more rapid than the rate-determining step, supporting the interpretation derived from examination of the early structures, which led us to focus instead on specific interactions between the ISP and its reaction partners that may restrict mobility by binding at the reaction interfaces. As we have shown, some of these can be measured directly through thermodynamic effects, and have provided the basis for our models for formation of ES- and EP-complexes [[4,34–36,40]. Some can be observed either through crystallographic solution [28,33,36,41,42], or through spectroscopic studies [42,43]. For many of these complexes, the dominant interaction, either observed directly or proposed, is through H-bonding between His-161 (bovine numbering is indicated by italics) of ISP and the occupant of the Qo-site. The nature of these interactions is strongly determined by the protolytic properties of this group, which are also reflected in the redox dependences of the half-cell reactions, and the pH dependence of redox titration data [40,44,45]. We have therefore invested much effort in characterizing these properties, and investigating how they are determined at the atomic level [43,46,47]. Our general approach in this latter investigation has been through specific mutagenesis, thermodynamic measurement (including protein film voltammetry, Table 1), crystallographic solution at high resolution, and detailed spectroscopic investigation of the local protein and solvent environment using pulsed-EPR, Resonance Raman (Fig. 2), and FTIR. Structures at high resolution (<1.5 Å) have been published for wild-type (2NUK) and 5 mutants (2NUM (Y156F), 2NWF (Y156W), 2NVE (S154T), 2NVF (S154C), and 2NVG (S154A) [47]. An additional structure for G133S has been described briefly in the context of spectroscopic data [46]. Five additional structures are at different stages of refinement for mutant strains modified at Leu-132 and Gly-133 (in collaboration with Satish Nair). Our spectroscopic studies have been published in four papers characterizing the WT protein in situ in the complex, and as the purified extrinsic domain [43,46,48,49]. Additional work is nearing completion, in which we have dissected the proton environment of the [2Fe-2S] cluster, and identified sites undergoing deuterium exchange, likely involved in H-bonds, and we have similar data from two mutant strains, Y156F and S154A, in each of which we have identified the loss of at least one exchangeable proton, with different properties in the two mutants. From the two structures for mutant strains at these different sites already published, we can say with confidence that the structures are essentially unchanged except for the loss of a single H-bond resulting from the mutation. It follows that the changes in thermodynamic and kinetic properties must arise from this change. For both mutations (Y156F, S154A), DFT/electrostatic calculations have recently been performed [50], and the results were in good agreement with our experimental data (Table 1), and support previous speculation [51–53] as to the importance of these interactions. From this extensive work we now have a detailed picture of the structural underpinnings of the thermodynamic properties of the ISP, and a much clearer understanding of how these properties are modified by mutation. At the resolution provided by our structures [46,47], crystallography provides true atomic-level positions for all non-H atoms, and in some cases also shows electron density for H-atoms. For some H-bonds, this makes it possible to identify angular parameters. This information is also in principle available from pulsed-EPR data. The special interest of our studies lies in the fact that we are accumulating such data from the same set of proteins, which will allow a crossvalidation of methodologies. Our recent observation that the backbone -NH of Leu-132 provides a H-bond to S2 of the cluster, and that the geometry leads to significant spin density on the N-atom (and by extension, on the S-atom of the cluster) is a nice demonstration of the synergy between these approaches [46].

1.2. Q_o-site reaction

From our earlier studies relating Berry's first complete structures to function, we had suggested a scenario in which movement of the extrinsic domain of the ISP was a spontaneous diffusional process with an expected time scale in the sub-microsecond range [28,35,36]. We demonstrated the feasibility of this scenario through molecular dynamic simulations. We pointed out that binding processes at the catalytic interfaces were likely relatively weak, based on the previously measured kinetic constants, and expected collision-limited rates, and on the occupancy of ISP in structures. In a more recent extended series of kinetic measurements [37], we were able to confirm these expectations, and demonstrate limits on reaction rates which were in line with these earlier speculations. These measurements provided a number of useful constraints on the mechanism.

a) We demonstrated that the oxidation of QH_2 in the absence of antimycin, determined from the electrogenic events associated with turnover, occurred at the same rate as that in the presence of antimycin, determined from the rate of reduction of heme b_H [37]. This result was important because it precluded any mechanisms in which physicochemical properties affecting turnover at the Q_0 -site were modified by binding of antimycin at the Q_i -site (cf. [9,10,53]).

b) We analyzed the rates of partial processes in the high-potential chain, and compare these with the turnover of the Q_o -site as determined by the kinetics of reduction of heme b_H in the presence of antimycin. We demonstrated that the 120 µs lag before onset of heme b_H reduction was almost all accounted for by the time needed to get an oxidizing equivalent to the Q_o -site. This result was important because it placed constraints on the movement of ISP, and on the stoichiometry of intermediate states, including that of the SQ. Since the rate-limiting step

Table 1	
Electrochemical parameters measured by CD spectrum,	protein film volatammetry, and kinetic titration

Strains	<i>E</i> _m values from CD spectra (mV)	Protein film v	Protein film voltammetry					Kinetic titration at 83% reduced quinine pool ^d		
		E_{acid} (mV)	E_{base} (mV)	pK _{ox1}	pK _{ox2}	pK _{red1,2}	pK _A	р <i>К</i> в	Vopt	
Wild-type/BH6 ^a	303±4 ^b /312±5 ^c	308±3	-134±6	7.6±0.1	9.6±0.1	12.4±0.4	6.66±0.10	9.14±0.26	973.3	
S154T	260 ± 5^{b}	288±3	-151±7	7.71±0.14	9.32±0.15	12.2 ± 0.1	6.83±0.12	8.48 ± 0.14	826.1	
S154C	313±5 ^b	324±4	-97 ± 6	7.17±0.11	9.77±0.13	12.0 ± 0.1	6.29±0.05	8.46±0.06	804.0	
S154A	167±6 ^b	178±4	-239±10	8.06±0.15	9.89±0.17	12.5 ± 0.1	7.52 ± 0.20	10.2 ± 0.28	199.2	
Y156F	256±4 ^c	252±4	-165 ± 6	7.76±0.15	8.83±0.21	11.8 ± 0.1	6.90 ± 0.06	8.54±0.08	714.6	
Y156W	198±3 ^c	208±3	-228 ± 4	7.87±0.09	9.25±0.11	12.2±0.1	6.97 ± 0.08	9.69±0.12	242.7	

^a The wild-type strain in this work is a complementary mutant strain of *Rb. sphaeroides* BC17, which contains the pRK415 with a modified *fbc* operon which has a histidine-tag at the C-terminus of the cyt *b* gene. The protein film voltammetric data of the wild-type was reported [44].

^b The values are measured at pH 7.4.

^c The data represent the limiting value for E_m at pH << p K_1 , as described in [59].

^d V_{opt} is the optimal reaction rate achieved when the first protonable group is deprotonated (pK_A) and the second group is protonated (pK_B), according to the following equation from the reference [148]. $v = V_{opt} \cdot \left(\frac{K_A}{K_A + |H^+|}\right) \left(\frac{|H^+|}{K_A + |H^+|}\right)$.



Fig. 2. Synergy of spectroscopy, crystallography, and thermodynamic analysis. (Left) Structures of wild-type, S154A and Y156F mutants (modified residues shown with electron densities); (middle) 2D ESEEM plots showing proton profiles in wild-type and S154A, and difference (bottom), showing loss of protons a, b (similar data have been obtained for the Y156F mutant); (right) RR spectra showing differences in ISP mutants S154A and Y156F, compared to SDX mutants and wild-type, and (bottom) PFV titrations, showing voltammogram (left, WT, pH 7), and *E*_m vs. pH curves from titration data for wild-type and five mutant strains (right).

was identified as the first electron transfer, with a rate constant of ~770 μ s, the unaccounted lag of <20 μ s would limit any occupancy to <0.026 for any intermediate between QH₂ and heme $b_{\rm H}$ [37]. This constraint on occupancy can be lowered to <0.015 on the basis of work from Millett's lab in which the reaction was initiated in the isolated *Rb. sphaeroides* complex by flash activation via a bound ruthenium complex [39,54]. This result eliminates all reaction mechanisms in which a stable (*K*>0.015) SQ-ISPH intermediate is required [55,56].

c) For the Q_o-site reaction itself, the first electron transfer (to ISP_{ox}) had been identified as the rate-determining step in QH₂ oxidation on the basis of dependence on driving force [25,26,57-59]. We had proposed a specific role for the H-bond between His-161 of ISP, and either QH₂ as H-bond donor in formation of the ES-complex, or Q as H-bond acceptor in formation of the EP-complex [25,40,45,59,60]. The difference between these roles depended on the protolytic properties of the histidine. We suggested that this group was responsible for the pK at 7.6 measured on the oxidized ISP. This pK was shifted to >12 in the reduced form. From this hypothesis, several predictions could be made about the properties of the first electron transfer reaction. The predictions were framed in the context of a mechanism in which the transfer of an electron (together with a proton) from QH₂ to ISPox occurred in a strongly endergonic process to generate the intermediate SQ at low occupancy. From the structures, and our proposal for the nature of the ES-complex, several paradoxical features had to be explained. These related to the high activation barrier, the slow rate of the overall reaction, and the very short distance for electron transfer (~7 Å). Since, after reduction, the ISP would be protonated, the reaction effectively involves H-transfer in a protoncoupled electron transfer. In the treatment proposed [4,40,42], the pK of His-161 contributed two separate effects. The first of these involved the role of the dissociated form of ISP_{ox} in formation of the ES-complex, in which QH₂ was the H-bond donor in the complex with ISP_{ox}. This meant that the concentration of ISP_{ox} as a substrate was determined by the pK on His-161. We demonstrated that this was the case by measurement of the pH dependence of electron transfer, which showed Michaelis-Menten saturation for ISPox [25]. Support for this interpretation was provided by use of a mutant ISP (Y156W) with a shifted pK, which also showed a corresponding shift in pH dependence [59]. The second effect of the pK on ISP_{ox} came from its role in determining the energy level of an intermediate proton configuration [61]. We suggested that electron transfer proceeded from an unfavorable configuration of the H⁺ in the H-bond, determined by a Brønsted relationship and the pK values for QH₂ and ISP_{ox}. The reaction interface was anhydrous, so this necessitated a protoncoupled electron transfer along the H-bond. We pointed out that the unfavorable intermediate state would contribute part of the energy barrier determining the rate of electron transfer, and that this could explain the observed properties. A detailed Marcus-type analysis showed that the behavior of WT could be accounted for economically by this mechanism, and the same treatment also explained the behavior of a set of mutant strains with modified $E_{\rm m}$ and pK values for ISP [4].

Because of the important role of the pK on *His-161*, and the coupling between redox and protolytic properties, we have invested much effort in determining the structural basis of these properties (see above). The attribution of the pK to a liganding histidine has now been confirmed by spectroscopy [62–64] and computational modeling [50] (see above). The mechanism proposed successfully accounts for the properties of the first electron transfer. Since this is the rate-determining step, our detailed mechanism in the context of the modified Q-cycle also explains much of the subsequent behavior of the complex.

d) We had proposed that the second electron transfer at the Q_0 -site might require a movement of the SQ from a location in the site distal from heme $b_{\rm L}$ where it was generated on oxidation of QH₂ to a proximal location, allowing ~1000-fold enhancement of rate constant and rapid reduction of the heme at low occupancy [25,60]. We had suggested a role for the conserved glutamate (Glu-295 of the -PEWYspan), which acted as a second H-bonding residue involved in formation of the ES-complex [34]. In this mechanism, oxidation of QH₂ by H-transfer to ISP_{ox} would generate the QH[•] form of SQ. Glu-295 would facilitated H⁺ released by accepting a H⁺ from OH[•], and rotating to deliver the H⁺ to a water chain leading to the exterior, leaving Q⁻ as the mobile form. Hunte et al. [8,65] have adopted a similar mechanism based on their yeast bc_1 complex structure, which was the first to show the water chain we had suggested from MD simulations [36]. More recently, Osyczka et al. [1,7] discussed possible mechanisms for the prevention of bypass reactions. They concluded that none of those previously proposed would work, because any configuration of the site allowing occupancy by SQ under circumstances in which heme b_L was reduced would give bypass rates similar to forward rates. We demonstrated that the alternatives discussed by them where physicochemically unrealistic, but that our earlier mechanism could function both to allow rapid forward electron transfer, and to limit bypass reactions by a coulombic gating [3,5]. We showed that the behavior of strains in which Glu-295 was mutated were consistent with the mechanism proposed. However, we should note that several other groups have constructed similar mutations in other systems, and although the results are for the most part consistent between groups, the interpretations have provided a different gloss on mechanistic aspects 66-68]. This residue is not absolutely conserved, and none of the mutants is completely inhibited, and some (for example E295D) show rates as high as 20% wild-type, and these facets have been interpreted as showing that Glu-295 is not mechanistically important. Our own analysis of bypass reactions clarified the arguments of Osyczka et al. [1,7], rejected their main conclusions, and provided severe constraints on possible mechanisms. The coulombic gating mechanism proposed involving Glu-295 offered a novel and successful resolution to the paradox they uncovered. A different gating mechanism was suggested by Mulkijanian [9], with alternative proton exit pathways, but a similar role for the glutamate. A role for glutamate in proton exit has been supported by the results of electrostatic calculations of different conformers at the Qo-site [69]. However, the site involves a metastable ES-complex, and because transitory states were not included in the calculations (which considered only the fully reduced or oxidized complex), the results can only be taken as providing an indication of potential protonation changes coupled to thermodynamically accessible states.

The question of the role of SQ in the Q_o-site reaction has been controversial for many years. Suggested mechanisms have ranged widely, from expectations for a major fraction of centers occupied by SQ in the intermediate state [31,55], through low occupancy models [25,26], to scenarios in which no SQ was expected [1,7]. In addition to the properties that led us to favor a low occupancy model [4,5,25], two other lines of evidence have strongly supported our suggestion that the SQ intermediate is formed at low occupancy by an endergonic proton-coupled electron transfer. Forquer et al. [70], in studies of by-pass reactions and super oxide (SO) formation, have confirmed that in yeast, as in the *Rb. sphaeroides* complex, the rate of the first electron

transfer is determined by driving force (and is therefore the limiting process), and have shown that the normal forward reaction, and the formation of SO both have the same activation energy. Secondly, Cape et al. [71] have demonstrated that SQ is formed at the site at low occupancy (~0.01) under anaerobic conditions in the presence of antimycin. Zhang et al. [72] have also demonstrated a SQ signal with a different line shape, also under anaerobic conditions, in a mutant truncated so as to remove heme $b_{\rm H}$, so that in each case electron transfer out of the *b*-heme chain was prevented. Each of these observations is consistent only with the second of these models. Cape et al. also showed that the SQ is not in H-bonding contact with any N-atoms (excluding a complex with ISP), but was in a constrained environment (assumed to be the Qo-site volume), as expected from the Hong et al. [25] mechanism. The occupancy of SQ was at the high end of the range expected from the rate of bypass reactions, but the occupancy expected was based on an assumed reaction distance similar to that in the ES-complex. The appearance of SQ was not time resolved in either of these studies. However, it is clear from the very low rate of bypass reactions under normal forward electron transfer that the re-oxidation of SQ keeps the level much lower than this [5], and it seems probable that SO appearance in these experiments required reduction of the *b*-heme chain by one or more turnovers of the Q₀-site before SQ could accumulate. This is likely also the explanation for the discrepancy between the results above [71,72] and the observation of Zhu et al. [73] in which no SQ was observed on initiation of turnover of the oxidized bovine complex by QH₂ addition. The authors make no mention of redox conditions, but the complex was initially fully oxidized and no antimycin was added, and it seems likely that the conditions were aerobic. Under aerobic conditions Cape et al. did not see SQ, in line with previous reports [26,74], and attributed this to reaction of SQ with O₂. The failure of Zhu et al. [73] to observe SQ could therefore be attributed to either its oxidation by O_{2} , or rapid re-oxidation by heme b_{1} , and is as expected from all these prior reports. Zhu et al. [73] reported the kinetics of reduction of heme $b_{\rm L}$ and ISP, and also showed raw data indicating the reduction of heme $b_{\rm H}$ and the appearance of SQ at the Q_i-site. For ISP and heme $b_{\rm L}$, 10% of the signal showed matched rates of reduction with $t_{1/2}$ ~200 µs. This half-time might imply a rate ~4-fold faster than the fastest rates previously reported in native photosynthetic membranes [25], or in the isolated complex [54], from Rb. sphaeroides, and ~25-fold faster than rates of Qo-site turnover measured by stopped-flow in the isolated bovine complex in the presence of antimycin. Taken at face value, the rapid and matched rates of reduction of ISP and heme $b_{\rm I}$ might be thought to place more severe constraints on the possible occupancy of SQ at the Q_0 -site than those previously suggested [5], but the noise level apparent in the EPR data shown would preclude more quantitative estimation. One possible explanation for the initial kinetics seen by Zhu et al. [73] is that the initial reaction represents a reduction kinetics truncated by re-oxidation, since the complex was initially fully oxidized. Indeed, extrapolation of the initial rate shows 50% completion at 1.1 ms, suggesting that the initial rate was in line with that measured previously for the rate-limiting first electron transfer reaction. If this is the case, and if a mechanism limited by the first electron transfer is assumed, the new data provide no tighter constraints than those previously estimated (see discussion above and [5,26,39,54]). The lower range of plausible occupancies calculated was $\sim 10^{-8}$ SQ/site [3–5,25], and such levels are certainly not precluded by the Zhu et al. [73] results. Any estimate of occupancy less than the previously calculated upper limit would make it more imperative to account for the rapid overall rate observed in terms of a rate constant for the second electron transfer faster than that possible if SQ had to react from the distal domain of the Qo-site [5]. No kinetic data in the presence of antimycin were reported by Zhu et al. [73], although effects of antimycin addition were discussed. De Vries et al. [75] had shown a faster rate of cyt $b_{\rm H}$ reduction in the absence than in the presence of antimycin, which was insensitive to myxothiazol, but the

data reported by Zhu et al. suggest a delayed reduction of heme $b_{\rm H}$ and a simultaneous appearance of SQ in the absence of antimycin.

1.3. Mechanism of the Q_i-site

Any mechanism must account for the fact that the Q_i -site operates as an interface between the 1-electron chemistry of the *b*-heme chain, and the 2-electron chemistry of the quinone pool, and involves intermediate states tuned to binding of the three components of quinone redox chemistry, Q, SQ and QH₂. Since the liganding requirements for these differ markedly, the site must be reconfigured during the catalytic cycle to accommodate these different requirements. In addition, pathways for uptake of two protons and for exchange of substrate and product need to be identified. It is generally supposed that the site operates through a two-electron gate in which heme b_H reduces Q to SQ on one turnover of the Q_o-site, and SQ to QH₂ on a second turnover [12,24].

1.3.1. The two-electron gate at the Q_i-site

The reaction at the Q_i-site has been treated as a simple twoelectron gate [24]. However, it has long been recognized that even such a simple treatment has built-in complexities relating to the involvement of three different components, O, SO and OH₂, as binding partners. The site has to accommodate each of these in any catalytic cycle, and each has its own requirements in terms of H-bonding [76]. The paradigm for discussion favored in the literature is in terms of the disproportionation reaction [77,78]. Values for the differential binding of Q and QH₂ and SQ stability are based on analysis of titration data from the EPR measurement of SQ. However, a disproportionation is clearly not appropriate as a description of the catalytic cycle, because, on the time scale of turnover, the site can only accommodate one species at a time, and the SQ is not exchangeable. Alternative approaches have been suggested, which attempt to reconcile the SQ data with the properties of the high-potential form of heme $b_{\rm H}$ associated with the cyt b_{150} phenomenon [79–81]. Our recent efforts on the Qi-site have been directed at the construction and characterization of mutant strains, and exploration of the local protein and solvent environment of the SQ form, since this information is accessible though pulsed-EPR [76,82-85. These have provided a detailed picture of the local structure [76,84], which, together with the available crystallographic data [41,65,86,87], has allowed new insights to mechanism. The pattern of H-bonding interpreted from crystallographic data has varied between labs. Berry's structures of the bovine and chicken mitochondrial complexes have shown a conserved His-201 and Asp-228 as ligands to carbonyl O-atoms of a bound quinone species, and the most recent structures of the Rb. sphaeroides complex from Xia's group, have also shown a direct ligand from the equivalent His-217. In structures of the yeast mitochondrial complex from the Hunte group, the H-bond to the histidine has been through a water molecule. In structures of mitochondrial complexes from Xia and collaborators, both H-bonds are modeled as through waters. Several of these structures are at close to atomic resolution, but because the quinone present can become reduced to SQ by X-ray exposure [88,89], the nature of the occupant in these structures is ambiguous. Our own EPR results are interpreted as showing ligands equivalent to those seen in Berry's structures (His-217 and Asp-252). However, MacMillan (personal communication) has reported that the N_{ϵ} of histidine seen in our ESEEM spectra is not visible in his using the yeast complex. The difference may come from the protocol used for generation of the SQ, - reduction by ascorbate at pH 7.8-9.0 in our experiments, and by dithionite at pH 9.5 in MacMillan's. The discrepancies between labs might also provide clues about variability in liganding of the three species occupying the site during catalytic turnover, and variability with pH on dissociation of liganding sidechains [76,84], but we have not yet explored all these possibilities. Our conclusions are summarized in a scheme in which conformational

plasticity allows the site to reconfigure to adapt to the binding requirements of the bound Q, SQ, and QH₂ forms that participate in the reaction [76]. It is likely that the primary ligands, His-217 and Asp-252, act as proton exchange sites; His-217 lies at the surface, so it provides a direct pathway, but Asp-252 is quite buried. Electrostatic calculations on the yeast complex suggest a strong coupling between residues equivalent to Asp-252 and Lys-251, other nearby lysines in cyt c_1 , and possibly bound cardiolipin. These are suggested to provide a channel for proton exchange with the aqueous phase [69].

Our more recent efforts have involved experiments to explore the N-environment, and the possibility of a third H-bond from Asn-221 (equivalent to Ser-205) [90]. In our previous work, no spin density on any additional N-atom apart from the histidine N_{ϵ} could be detected using ¹⁴N X-band. From the ¹⁴N S-band and ¹⁵N X-band spectra, we have now determined the complete nuclear quadrupole tensor of the ¹⁴N, and the isotropic and anisotropic couplings, with the N_e of His-217. However, no significant spin density on other N-atoms was detected. DFT calculations showed that the energies expected from the Asn-221 H-bond would be masked by the dipole-dipole signals from other N-atoms, leaving the question open. Several mutant strains at Asn-221 were constructed, were able to grow photosynthetically, and showed turnover of the Qi-site reactions, but with inhibition in rate, and modified properties for heme $b_{\rm H}$. We therefore concluded that the potential H-bond from Asn-221 is not essential, but is important for function. The characteristics of the weak H-bond observed through exchange with deuterium suggest that it involves a bound H₂O, but some of the diagonal peaks in ¹⁵N HYSCORE spectra could have been contributed by a H-bond from Asn-221. The most recent structures [87] include one showing a quinone occupant (PDB ID 2qjy), in which N_{ε} of His-217 serves as a direct H-bonding ligand, as seen by EPR [90], no H-bond from Asn-221, and a distance to Asp-252 too great for a direct H-bond, but compatible with a watermediated H-bond. The structure most likely reflects the occupancy by the oxidized Q.

1.3.2. The role of cyt b_{150} in the mechanism of the Q_i -site

The component known as "cyt b_{150} " is due to a perturbation of the properties of a fraction of heme $b_{\rm H}$ on interaction with SQ at the Q_i-site. It is seen in redox titrations of chromatophores [91], mitochondria [92] or isolated complexes [93], as a component titrating with $E_{\rm m,7}$ 150 mV only in the absence of antimycin [81]. From the properties of cyt b_{150} we suggested that the Q_i-site catalyzes the following reaction, in which the cyt b_{150} is represented by the state cyt $b_{\rm H}$ HQ⁻⁻ [2,94]. The reaction (Eq. (1)) can be considered as a reversal of the normal forward reaction by which ferroheme $b_{\rm L}$ reduces SQ to QH₂, and, after release of the product, binds Q to re-establish the starting state for the Q_i-site.

$$QH_2(pool) + cyt \, b_H Q \leftrightarrows Q(pool) + cyt \, b_H H \, Q^{-}(H^+)$$
(1)

This overall reaction can be split into partial processes:

$$QH_2(pool) + cyt \, b_H Q \leftrightarrows Q(pool) + cyt \, b_H QH_2$$
(1a)

$$\operatorname{cyt} b_{\mathrm{H}} \mathrm{QH}_2 \leftrightarrows \operatorname{cyt} b_{\mathrm{H}} \mathrm{H} \mathrm{Q}^{-}(\mathrm{H}^+) \tag{1b}$$

allowing definition at any defined pH, of an equilibrium constant for formation of the cyt b_{150} form:

$$K_{eq} = K_{(1a)}K_{(1b)} = (K_{A(QH2)}/K_{A(Q)})exp((E_{m(cytbH)}-E_{m(QH\cdot/QH2)})F/RT))$$
(2)

where K_A values are association (binding) constants for QH₂ and Q, and $E_{m(cytbH)}$ and $E_{m(QH^{-}/QH^{2})}$ refer to the mid-point potentials of cyt b_{H} , and of the SQ/quinol couple bound at the site, respectively.

The affinities for Q and QH_2 can be subsumed under the terms for the potential of the quinone pool and the Q/SQ couple at the site, since the ratio of binding constants for Q and QH₂ contributes to the value for $E_{m(QH+/QH2)}$ so as to cancel the ratio of binding constants. This gives:

$$K_{eq} = \exp((E_{m(cytbH)} + E_{m(Q/QH)}) - 2E_{m(Qpool)})F/RT$$
(3)

(where $E_{m(Q/QHP)}$ refers to the bound couple), which can also be derived directly from Eq. (1). A computer model based on this set of equations allows exploration of the variation of redox states of all components as a function of redox potential, pH, etc. In this treatment, no account is taken of different protonation states, local coulombic effects, etc., and these effects are subsumed under existing terms. However, the observed pH dependence of both the b_{150} and SQ titrations are handled by a single pK value, currently assigned to SQ (see discussion below).

The mechanism proposed here is in contrast to the mainstream of discussion which starts from consideration of the disproportionation reaction as a basis for mechanism [77.82.95.96]. Since structures became available, there has been little discussion of how this might occur at a site that appears too small to contain more than one species. The Qi-site catalyzes a quinol-quinone transhydrogenase reaction that allows transfer from exogenous quinol to bound quinone, or between exogenous quinones [97,98]. A plausible mechanism would be for two electrons from a donor Q_dH_2 to be stored in the two *b*-hemes, allowing dissociation of the Q_d, and its replacement by an acceptor Q_a, which would then leave the site as $Q_a H_2$ [98]. The rate of such a reaction would depend strongly on the probability of populating the $b_{\rm H}^{-}b_{\rm L}^{-}$ state. The unfavorable equilibrium constant for reduction of heme $b_{\rm L}$ through b_H [12,99] would account for the relatively slow rate observed. Nevertheless, this would likely be sufficient to allow equilibration on the time scale of a redox titration. Equilibration through 1-electron transfer processes might also occur through direct or indirect interaction with mediators. Since the conditions used to assay semiguinone by EPR (high concentrations of redox mediators, and a long equilibration time) would allow the reaction to reach equilibrium, it is reasonable to use the data in discussion of mechanism, but it is highly unlikely that it reflects the kinetically important intermediate states.

Since the hemes are 1-electron acceptors, it is necessary to take account of the equilibration of the Q/QH_2 couple with heme b_H through 1-electron transfer reactions. — an attractive feature of the cvt b_{150} mechanism proposed. Reduction of heme $b_{\rm H}$ through the Q_i-site by exogenous quinones, with generation of a semiquinone, occurs on addition of quinol substrate to the oxidized complex in the presence of myxothiazol. In a definitive study by de Vries [75], reduction by duroquinol led to generation of ferroheme $b_{\rm H}$ and SQ simultaneously, with the reaction ~90% complete at the first point of measurement at 5 ms (using 300 μ M QH₂). From this, a value of τ <3 ms seems appropriate. In chromatophores, where the reactions using the native substrate can be followed in situ, the myxothiazol insensitive reduction of heme $b_{\rm H}$ on generation of 1 QH₂ in the oxidized pool also occurred with τ ~3 ms [80]. Since this is ~two orders of magnitude faster than the transhydrogenase reaction, the disproportion reaction is excluded. It seems likely that duroquinol, and presumably other quinols, can react directly and rapidly at the Qi-site by exchange with the initial occupant.

Rich et al. [96] have discussed the cyt b_{150} phenomena in terms of differential affects of occupancy by Q, SQ or QH₂ on the E_m of heme b_H (see also [82]). A set of 7 different E_m values and 2 pKs was required to account for the properties of both the SQ and cyt b_{150} , but these parameters did not include the possibility for differential binding of the different components. Furthermore, in deriving this set, Rich et al. [96] had to assume that SQ generated in the presence of oxidized heme b_H was EPR silent due to magnetic quenching (see also [100]). If the semiquinone signal at the Q_i-site is modulated by magnetic interaction with the spin of oxidized *b*-heme [100,101], this would lead not only to loss of amplitude, but also to some displacement of the maximum of the bell-shaped titration curve [96], and hence to misleading values for

derived thermodynamic parameters. There is no doubt that occupancy by inhibitors can modify properties of heme $b_{\rm H}$, and a substantial literature showing changes in both $E_{\rm m}$ and $\lambda_{\rm max}$, but the complexity of the Rich et al. treatment makes it somewhat intractable to testing. In contrast, the model proposed here has only a single equilibrium constant, and singular $E_{\rm m}$ values for heme $b_{\rm H}$ and the quinone system taken directly from experiment. Computational modeling shows that the properties of the cyt b_{150} phenomena are reproduced quite well using measured values for $E_{\rm m}$ for heme $b_{\rm H}$, values for binding constants and E_m values for the Q/SQ/QH₂ system derived from redox titrations and kinetic experiments. The properties of the SQ are also reproduced quite well, but are mechanistically linked to formation of SQ through reaction 1 above, so that the SQ observed is in the form cyt b_HH'Q⁻. An interesting feature of these simulations was that, because of the importance of $E_{\rm m}$ of heme $b_{\rm H}$ and the Q/SQ couple in Eq. (3), the pH dependence of both b_{150} and the SQ were well explained, but are determined by a single pK. Previously we have used the published pK at ~7.8 on the heme $b_{\rm H}$. However, we have recently revisited these titrations (see below) and now prefer a pK in this range on the semiguinone, or associated with interaction with the protein (cf. [69]).

1.3.3. The forward reaction

The mechanism for formation of the cyt b_{150} state provides the main elements of a mechanism for the forward reaction [2]. Additional parameters needed are those for reduction of Q to SQ by electron transfer from heme $b_{\rm H}$. The fact that the observed rate is that of the limiting step at the Qo-site restricts use of the kinetic data to estimation of lower limits for the forward rate constants. However, the equilibrium constant can be estimated from the kinetics with the quinone pool initially oxidized (and the site presumably occupied by Q). After flash activation to populate the ferroheme $b_{\rm H}$ (generated as the one QH₂ in the pool produced by excitation of the RC is oxidized through the Qo-site), the heme remains partly reduced in the 100 ms range, suggesting an equilibrium constant for reduction of Q by heme $b_{\rm H}$ H with K_{eq} ~3. This relatively stable level of heme $b_{\rm H}$ reduction titrates away as the pool starts to become reduced over the range below $E_{h,7}$ 150 mV. This is the range over which the cyt b_{150} state is populated, so the acceptor at the site will change from Q to SQ over this range. A mechanism for the forward reaction can then be formulated, as discussed more extensively elsewhere [2].

2. Can a monomeric mechanism account for the observed functional data?

The structures all show a dimer, which raises the main question to be discussed in the remainder of this paper. What do we know of the consequences of this dimeric nature? Are there mechanistic parameters dependent on interactions across the dimer interface?

The structures now available (including several structures from two Rhodobacter species [31,32,87]) have demonstrated the high degree of conservation of catalytic subunits across the bacterial/ mitochondrial divide, highlighted the dimeric nature of the complex, and provided a framework for extensive discussion of the role of the dimer in mechanism [6,7,53,102-106]. The attractiveness of the modified Q-cycle treatment rests on the fact that the properties can be related to a few constants, measured directly from thermodynamic or kinetic experiments on the intact membrane systems, or calculated from structural data. It is appropriate to emphasize that the kinetic data for native and mutant strains accumulated over the past 25 years have allowed us to subject the modified Q-cycle model to detailed analysis in terms of stoichiometric ratios of reactants, the equilibrium constants determining their distribution, and the kinetic constants determining the time course of changes on introduction of substrates [12,37,107–110]. Because the system can be flash-activated, no mixing of substrates is needed, and as a consequence, the initial state of the

system can be manipulated over a wide range of $E_{\rm h}$ and pH to change substrate concentrations, thermodynamic poise, etc. These data are satisfactorily and economically explained by the monomeric modified Q-cycle. No ad hoc intrusions spoil the simplicity; - there is no need to introduce secondary interactions between sites, or actions-at-adistance, beyond those introduced by simple coulombic considerations. The mechanism is completely natural, - a conclusion reinforced by more recent data in which kinetic contributions were more accurately resolved by deconvolution, allowing finer points to be explored [111-113]. This is in contrast to schemes introduced by other groups, in which mechanistically occult interactions between catalytic centers, or between reaction intermediates and redox states, have been proposed, which have been given prominent controlling roles [10,53,103,104]. These speculative interactions are based on interpretation of kinetic experiments, usually using an isolated mitochondrial complex in stopped-flow mixing protocol. Although this approach has been applied with skill, intrinsic difficulties limit the utility of some experiments. These arise from the nature of the system, - the experiments require the detergent solubilized complex, and use of more soluble guinol substrates than the native ubihydroguinone-10. As a result, the binding of QH₂ leading to formation of the ES-complex is always unnatural. Even with decylUQH₂ or analogs with truncated isoprenyl sidechains, binding involves partition from a mixed aqueous and micellar phase in which the guinol has a limited aqueous solubility [114] (CMC~30 µM). Often, quinones without the lipophilic side chain, and with very different redox properties (menaquinone or duroquinone, for example) have been used, and these differences change both the kinetics of binding and the equilibrium constants for partial processes, with sometimes dramatic kinetic consequences ([115] and Kokhan, O. and Wraight, C.A. personal communication), so that the observed effects often differ from those seen *in situ*. Because of the limitations of stopped-flow, one has to be careful in interpreting the results, - in particular when using them to justify hypotheses about the physiological mechanism. There is a strong temptation to presume that any kinetic effect that cannot be understood in terms of the simple model must reflect some new twist to the Q-cycle theme, and this has encouraged much speculation centered on "hidden" interactions. Unfortunately, the scope for speculation increases dramatically, with a power-law dependence on the number, *n*, of such interactions proposed, further compounded by a square function if interactions between monomers in the dimer are considered.

We will take as a starting point for discussion the mechanism summarized in Fig. 1, and examine the hypothesis that this minimal model of a monomeric mechanism [2–5] is sufficient to account for the metabolic role of the bc_1 complex. Although the structure is dimeric, the model can accommodate those features from the structural evidence that involve the dimer in a scaffolding role. The ironsulfur subunit of one monomer is structurally constrained by specific interactions at the membrane level, involving a clamp between cyt b subunits, and interactions with other subunits mainly in one monomer, which leave the extrinsic mobile head free to interact with catalytic interfaces in the other monomer. This dimeric involvement could in principle account for the dimeric nature without any extraneous functional role. However, several labs have claimed to demonstrate inter-monomer electron transfer [6,53,103], and others have assumed on the basis of strong theoretical grounds that this must occur [7]. Furthermore, a substantial body of evidence has shown that binding of antimycin modifies the interaction of ISP so as to change the susceptibility to proteolytic cleavage [116], and recent results from Daldal's group have shown conditions in which the g-tensor of the [2Fe-2S] cluster appeared to change with respect to the orientation of chromatophore films on binding of antimycin [102]. Both these latter effects are discussed in the context of a modified interaction with the Qo-site occupant, reflecting a coupled conformational change induced by antimycin binding at the Q_i-site.

As several commentaries have noted [7,27,53,117], the structures show distances between the b_1 hemes across the dimer interface that are expected to facilitate rapid electron transfer. These are Fe-Fe at 21.58 Å, nearest conjugate atoms at 10.54 Å (vinyl sidechains), and pyrrole rings at 14.45 Å (in 1pp9) (Fig. 3A). Values for rate constant depend on which distances is used, and on whether a classical Marcus [118,119] or Moser-Dutton [120-124] treatment is used for the exponential term. Some representative values for the competing pathways are shown in Table 2. If the appropriate distance is that between conjugate systems, it would likely include the vinyl sidechains, the path assumed in the Table. The values for ΔG chosen depend on the redox status, and hence on any effect of coulombic interactions between hemes. For example, if heme $b_{\rm H}$ is reduced in one monomer, the coulombic effect on b_L will be about -60 to -80 mV, generating an exergonic pathway for electron transfer to the neighboring heme $b_{\rm L}$, if the heme $b_{\rm H}$ in that monomer is oxidized (bottom line). Note however that all b_L - b_L rate constants would allow rates more rapid than the rate-determining step ($\sim 1.35 \times 10^3 \text{ s}^{-1}$) [25,26]. A succinct review of this problem by Shinkarev and Wraight [125] is couched in terms of competition between the heme $b_{L} \rightarrow$ heme $b_{\rm H}$, and heme $b_{\rm L1} \rightarrow$ heme $b_{\rm L2}$ paths. In assigning rate constants, they assumed the appropriate distance was that between conjugate rings, (14.45 Å), but since the vinyl groups are part of the conjugate system, it does not seem natural to exclude them, and this increases rate constants by a factor of ~200. Then, although competitive effects could lower expected rates as they suggest, under conditions in which both hemes $b_{\rm H}$ are oxidized, no combination of effects within the conventions of the Moser-Dutton treatment of the intrinsic rate constant would prevent electron transfer across the dimer interface at rates in the range of the limiting rate once the "active side" $b_{\rm H}$ became reduced. The expectation is therefore that electrons could be shared between monomers at this level, and that this pathway would be functional, and easily observable on the time scale of ~100 ms used in typical experiments.

Experimental evidence in support of dimeric operation has come from several studies. Covian et al. [126] interpreted stopped-flow kinetic experiments as showing that the stoichiometry of cyt c_1 reduction on addition of decylUOH₂ was half that expected from the total complement, and that only half of the complex present participated in electron transfer. Under these conditions, two cyt *b*-hemes became reduced per active enzyme in the presence of antimycin (although no effort was made to distinguish between hemes $b_{\rm H}$ and $b_{\rm L}$). These properties were similar to previously observed kinetic anomalies, which were explained by the modified Q-cycle. For example, on flash activation in the presence of antimycin, a discrepancy between amplitudes of heme b and heme c reduction is observed, and had been seen as contrary to the expectations of a Q-cycle [127]. However, the modified Q-cycle requires two turnovers of the Qo-site for a full turnover of the complex, and detailed analysis in the early 1980s showed the equilibrium constant for these depends on the acceptor. The value depends on $E_{\rm m}$ for heme $b_{\rm H}$ on the first turnover $(K_{eq} \sim 480)$, and that of b_L on the second turnover $(K_{eq} \sim 3)$, and the degree of reduction of the high-potential chain is simply a reflection of this change in the nature of the acceptor [12]. However, Covian et al. [126] explained their results in terms of an anti-cooperative mechanism in which the Qo-site of only one monomer was functional, but fed into both sides of the dimer through electron exchange at the heme $b_{\rm L}$ level. In support of this, they showed titration curves with antimycin of the initial rate of cyt c reduction in a steady-state assay of activity that gave a convex curve, with a stimulation of rate in the sub-stoichiometric range. In a later paper [6], the same group reported titrations by antimycin of cyt *b* reduction on addition of decylUQH₂ to the myxothiazol inhibited complex, in which the titration curves showed a convex shape. They again interpreted the results as showing electron transfer between the $b_{\rm L}$ hemes.

Although such convex curves might be expected from electronsharing, a substantial literature from the 1970s, following the observation of Kröger and Klingenberg [128,129] of hyperbolic titration curves



Fig. 3. Schemes to show potential electron transfer pathways involved in electron equilibration between monomers at the level of heme b_L . Top. The redox centers of the dimer are shown, and the Q_o - and Q_i -sites indicated by stigmatellin and antimycin, respectively. The scheme shows the dimer with the Q_i -site blocked in both monomers by antimycin (blue crosses), and the Q_o -site of one monomer blocked by myxothiazol (red cross). Under these conditions, if electron transfer between hemes b_L could occur, both hemes b_H could be reduced by turnover of the uninhibited Q_o -site (broken red arrows). Bottom. The dimer interface, showing residues referred to in the text. The dotted line shows the 10.21 Å distance between heme b_L 4-vinyl groups (taken from PDB file 2qjy, chains A–F).

with antimycin resulting from a "pool" effect of ubiquinone, demonstrates that alternative explanations are possible. In general, a Kröger– Klingenberg effect will be found whenever the turnover of the process being titrated is greater than that of the rate-limiting reaction, since a fraction of centers can be eliminated without a full effect on the overall process. We have been able to generate convex titration curves under a variety of such conditions with chromatophores ([130], and in collaboration with Vlad Shinkarev), or with isolated bc_1 complex. However, when care was taken to ensure that the rate-limiting step was the process titrated, linear curves were always seen. Indeed, such linear titrations are the norm in the literature. Several other mechanisms can give non-linear titration curves, as shown on titration of the amplitude of cyt c_1+c_2 oxidation following flash activation in chromatophores [130], where the degree of convexity is dependent on which time-point in the kinetic curve is chosen. This is because mobility of the substrates QH_2 and cyt c_2 allows one active complex per chromatophore to rereduce all cyt c_2 , given time. This effect has an *n*-value from Hill plots of ~9, likely reflecting the number of bc_1 complexes per chromatophore. Similarly, titration by antimycin of the kinetics of cyt b_H reduction and

Table 2

Reaction pairs and conditions	Moser–Dutton k/s^{-1}	Marcus k/s^{-1}
b_L-b_H 2-vinyl (assuming $\Delta G = -60 \text{ mV}$, $\lambda = 0.75$, $R = 7.02 \text{ Å}$)	8.97×10 ⁸	1.73×10 ⁸
$b_{\rm L}-b_{\rm H}$ 2-vinyl (assuming ΔG = -130 mV, λ = 0.75, R=7.02 Å)	2.15×10 ⁹	5.68×10^{8}
b_L-b_L 4-vinyl (assuming $\Delta G = -0$ mV, $\lambda = 0.75$, $R = 10.54$ Å)	2.86×10^{6}	4.08×10^{5}
b_L - b_L 4-vinyl (assuming ΔG =-60 mV, λ = 0.75, R=10.54 Å)	6.51×10^{6}	1.25×10^{6}

re-oxidation shows a similar effect, possibly, as suggested by Bechmann et al. [131], because antimycin can leave one Q_i-site and find another, so long as one remains open. In the isolated complex this gave an *n*-value

of 2, but we found a higher value in chromatophores (>4.9) (Fernandez-Velasco and Crofts, unpublished), which suggests that the inhibitor can redistribute as long as an open site is available in a chromatophore. In



Fig. 4. Titrations of Q_o -site with myxothiazol showing linear curves in WT and mutant strains. A) Typical curves showing the kinetics of heme b_H reduction in WT in the presence of antimycin. Similar sets of data were taken for each strain, all of which had similar maximal rates. Myxothiazol concentrations are indicated to the right of each curve. B) Fraction of heme b_H reduced at 20 ms (expressed as %) in the presence of Q_I -site inhibitor antimycin A, as a function of inhibitor concentration. The system was excited with a xenon flash (~5 µs at half-height) and the reduction of b_H monitored from the difference kinetics measured at 561–569 nm. Ambient redox potential was poised at ~100 mV (QH₂ oxidation close to maximal). Strains are indicated by labels; bc_1 complex in the range 70–220 nM.

both cases, linear titrations are seen under conditions in which the ratelimiting step was titrated.

$$K_{\rm eq} = \frac{k_{\rm f}}{k_{\rm b}} = \frac{[b_{\rm L}^-]^2}{[b_{\rm f}^-]^2} = 10^{\Delta E_{\rm m}/(59.19)} = 0.054$$
(4)

$$K_{\rm eq} = \frac{k_{\rm f}}{k_{\rm b}} = \frac{[b_{\rm H}b_{\rm L}^-]}{[b_{\rm H}^-b_{\rm L}]} = 10^{4E_{\rm m}/59.19} = 0.0029$$
(5)

In the work from Covian and et al., the case for an interpretation in terms of electron exchange between monomers depended on a kinetic model in which a questionable value for the equilibrium constant between hemes $b_{\rm H}$ and $b_{\rm L}$ was used [6,126]. The treatment involved a reaction equation, $b_{\rm H}^- + b_{\rm L} = b_{\rm H} + b_{\rm L}^-$, appropriate for the reaction in the solution. Then a condition $[b_{\rm H}^-]=[b_{\rm L}]$, and $[b_{\rm H}]=[b_{\rm L}^-]$ was imposed because the actual reaction was intramolecular (Eq. (4)), and this changed the equilibrium constant determining occupancy of the $b_{\rm I}$ state by 18.6-fold (compare Eqs. (4) and (5)). The value derived allowed sufficient occupancy of the $b_{\rm L}^-$ state to provide an inter-monomer rate that explained the observed rate. Without the larger value for K_{eq} their model fails to match the kinetic data. The treatment is in error on both mechanistic and thermodynamic counts; the reaction is not 2nd order, and by assigning separate identities to four reactants, they are trying to have their entropic cake and eat it too. Intramolecular electron transfer is an exchange between two states in which the electron is on either the one or the other center, and treating the system as 1st order gives the conventional equilibrium constant expected from a one-electron transfer. For the transition $b_H^- b_L \Rightarrow b_H b_L^-$, K_{eq} is determined as in Eq. (5). Any alternative path with a different equilibrium constant would make possible a perpetual motion machine. Because the experiments of Covian et al. [6,104,126] did not discriminate between several possible mechanisms, and because their model is suspect, it can hardly be taken as established from these results that the mechanism is dimeric, or that electron exchange must occur between monomers on the time scale of turnover

Gong et al. [104] have constructed mutants in Rb. sphaeroides with changes at three aromatic residues, Phe-195, Tyr-199, and Phe-203, which, on the basis of a computer model, were expected to line the dimer interface between the $b_{\rm L}$ hemes. They studied steady-state electron transfer and SO production in bc1 complex isolated from variants in which each residue was changed to alanine, with some additional changes at F195. The electron transfer rates reported (2.5 µmol cvt c reduced/nmol heme b/min or 21 mol/mol/s for the wild-type) were only ~1.5% physiological rates (1350 mol/mol/s at saturating substrate), indicating that the turnover was substantially inhibited in all these preparations. None of these mutations had a strong inhibitory effect. The greatest effect was seen when Phe-195 was changed; F195A had a lower rate (80% of WT) than F195W, F195H or F195Y, and a higher rate of SO production. On this basis, the authors concluded that an aromatic residue was required, and that the critical function was the electron transfer between the $b_{\rm L}$ hemes. However, this electron transfer path was conjectural; no evidence was presented showing that any such reaction occurred, and the effects could as well have been attributed to a weak inhibition of the Qo-site, which (on the basis of myxothiazol occupancy) is 14–19 Å from these residues. In structures now available, only Tyr-199 is in the most direct path, whether interpreted as through bond or through protein. We discuss below our own results based on a series of mutations at this residue, Y199T, S, C, W and L, and show that these are clearly against an inter-monomer exchange.

In another challenge to the modified Q-cycle, Mulkidjanian [9,10] has suggested an "activated" Q-cycle with a number of novel features, at least some of which seem untenable. He attributed the acceleration of the rate of QH₂ oxidation in the range $\sim E_h$ 150 mV to a priming process involving formation of SQ at the Q_i-site (the cyt b_{150} effect).

However, the same acceleration over this E_h range occurs in the presence or absence of antimycin, which specifically eliminates the Q_i -site SQ, so this must be wrong. It had previously been demonstrated that this acceleration was due to an increase in [QH₂] on reduction of the Q-pool, and preferential binding of QH₂ at the Q_o-site [12]. Another odd experimental result under special conditions was a marked difference in rates of turnover of the Q_o-site in the presence or absence of antimycin. However, this contradicts studies from a number of labs [12,37,132–135]] showing the rates to be the same under these two conditions. It seems likely that these aberrant observations reflect a lack of technical finesse rather than real functional differences.

Other groups have suggested control functions based on conformational coupling linked to structural changes on occupation of the Q_o -site by different redox species [39]. However, the causal relationship proposed could be as well explained by dynamic flexibility to accommodate different reaction partners [40].

3. Myxothiazol titrations to test for electron transfer between monomers

In the presence of antimycin, oxidation of heme $b_{\rm H}$ is prevented, so electrons passing out of the Q_o -site accumulate in heme b_H . In chromatophores, the stoichiometry of reaction centers to bc_1 complex is about 2:1, ensuring that sufficient substrates (2 cyt c_2^+ plus 10H₂) are generated on a saturating flash to allow one full turnover of the complex. In the presence of antimycin, the two oxidizing equivalents in the high-potential chain could in principle oxidize 2 QH₂, with reduction of hemes $b_{\rm H}$ and $b_{\rm L}$. However, the equilibrium constant for the second turnover, resulting from the low $E_{\rm m}$ (~-90 mV) of heme $b_{\rm L}$, prevents this, restricting the complex to 1 turnover of the Qo-site [24,109]. If inter-monomer transfer could occur [27,53,104,117], electrons from either Q₀-site in the dimer could reduce both hemes $b_{\rm H}$. Even with one monomer per dimer inhibited at the Q_o-site by myxothiazol, full reduction of heme $b_{\rm H}$ should be observed, because both $b_{\rm H}$ centers would operate at the same higher $E_{\rm m}$ (~40 mV), so that the equilibrium constant would strongly favor reduction. In practice, statistical effects would lead to a bowed titration curve (for example, at 50% occupancy, the distribution of dimers expected is 25% each of 0,0; 0,1; 1,0; and 1,1, where 0 and 1 are vacant and myxothiazol inhibited sites, so 75% of the full reduction might be expected). This convex titration curve is not seen. This could possibly be attributed to some special role for the intervening protein. We have constructed mutant strains in which the residue at the contact point between monomers at the level of heme b_1 , Tyr-199, was modified to give Y199 W, T, S, L, and C. Since this residue is next to His-298, a ligand to heme $b_{\rm I}$, and lies on the most direct path of electron transfer between hemes $b_{\rm I}$ (Fig. 3B), we anticipated substantial effects when inter-monomer electron transfer was probed by titration with myxothiazol (Fig. 4). Fig. 4 shows the results observed. All strains showed similar rates for Qo-site turnover in the absence of myxothiazol. None showed any marked change in redox properties of the cytochromes (Table 3), although all showed small changes in the α -band spectrum of heme $b_{\rm L}$ Titrations for all strains (including WT) were strictly linear, whether measured through initial rate, or amplitude of reduction, or

Table 3

Strain	Redox potentiometry <i>E</i> _m , (V)			Nernst equation solution fitted at each wavelength of the spectrum, E_{m} , (V)				
	b _L	$b_{\rm H}$	b ₁₅₀	Cyt c ₁	b _L	b _H	b_{150}	Cyt c ₁
WT+antimycin	-0.113	0.049	-	0.250	-0.090	0.040	-	0.250
WT	-0.120	0.054	0.169	0.250	-0.090	0.040	0.150	0.250
Y199L	-0.123	0.038	0.149	0.259	-0.100	0.030	0.148	0.250
Y199S	-0.077	0.057	0.172	0.265	-0.094	0.053	0.170	0.260
Y199T	-0.108	0.045	0.160	0.262	-0.092	0.027	0.152	0.250
199-insert	-0.096	0.035	0.155	0.251	-0.092	0.033	0.135	0.250
Y199W	-0.080	0.075	0.170	0.237	-0.091	0.050	0.150	0.250

after 1 flash or two (Fig. 4B). Since the second flash leads to re-oxidation of the high-potential chain, the driving force for inter-monomer electron transfer would be substantially increased, and an "extra" reduction of heme $b_{\rm H}$ would have been expected if electron transfer between the $b_{\rm L}$ hemes could occur, but this was not observed. Instead, heme $b_{\rm L}$ went reduced after the second flash, as expected in the monomeric Q-cycle [12]. Furthermore, no indication of the biphasic character expected from a branched pathway was apparent in the kinetic curves. We also tested these properties in a strain (H198-T199-Y200) constructed so as to mimic the extra residue in the sequence at this point found in the cytochrome $b_{\rm ef}$ complex of the cyanobacterial/ chloroplast photosynthetic chain (with sequence -HTF-) [136]. Again, myxothiazol titrations were linear, although the structures now available show a distinctive change in configuration of the interface in the $b_{G}f$ complex compared to the bc_1 complex, which might also have been induced by the mutation. The simplest interpretation of these results is that no inter-monomer electron transfer occurred, despite the short distance between the hemes and the rapid rate constant expected.

The above conclusion presents an interesting paradox. From treatments using the Moser–Dutton approximation for intrinsic rate constant at this distance, electron transfer should occur, but the experiments show that it does not. If unequivocal evidence becomes available showing that



Fig. 5. Coulombic free-energy change expected between hemes. The change in $E_{\rm m}$ expected from coulombic effects between hemes calculated from distance apart, for different intervening dielectric values.

electron-sharing occurs, then the data interpreted as showing that it does not will have to be re-evaluated. If on the other hand, no such evidence is found, then it will be necessary to explore the possibility that other parameters must be important in determining electron transfer rate, over and above those in the Moser–Dutton treatment. Either result would be of interest, and advance the field.

Several possibilities look interesting. The Moser-Dutton treatment uses the Hopfield approximation [119] to allow for quantum mechanical effects, but this effectively modifies $k_{\rm B}$, giving rise to thermodynamic inconsistency in applying the Marcus term to the Arrhenius expression [137]. From the Marcus curve generated using Moser–Dutton value of 3.1, values for $\log_{10}k$ at any chosen ΔG appropriate for a forward reaction, and for a reverse reaction at a value taken symmetrically with respect to $\Delta G=0$, do not give the ratio expected from the relation of ΔG to K_{eq} . In the Moser–Dutton treatment of the intrinsic rate constant, the approach is implicitly classical, since fluxes through all pathways are additive, and distance and density are taken to provide all information pertinent to the probability of electron transfer through the protein matrix. Alternative approaches have treated the intrinsic rate constant quantum mechanically [119,138]; then a standard path integral treatment opens the possibility of destructive interference, so that, in this scenario, interference in the inter-heme $b_{\rm I}$ path could limit rate. Indeed, several recent papers [139-143] have considered more sophisticated treatments that include interference terms. In a few cases, especially those involving pathways including heme and the liganding histidine, destructive interference was found to play an important role. Additional complications arise in systems involving electron transfer between subunits [142]. A more conventional approach is in terms of the electron distribution on the hemes. If the electron distribution at the 2-vinyl position is more favorable than at the 4-vinyl, this would favor electron transfer from heme $b_{\rm L}$ to $b_{\rm H}$. Walker [144,145] has kindly provided useful insights into the electronic structure as determined by calculation and EPR, and has provided data showing that the β -carbons of the 2-vinyls of the ferrihemes should both have substantially higher spin occupancies than those of the 4-vinyls (Ann Walker, personal communication). Since rate depends on *k*[occupancy], the $b_L \rightarrow b_H$ pathway (in which the 2-vinyls are in the shortest path) should be favored.

4. Coulombic interactions in the dimer

When the bc_1 complex was mutated to replace one of the histidine ligands to heme b_{H} , the complex was still assembled with a full



Fig. 6. Thermodynamic behavior of the *b*-type hemes in H6B strain (his-tagged cyt *b* with wild-type sequence) *in situ*. (Top left) typical titration data in the absence and presence of antimycin (pH 7); (bottom left) spectra from redox cuts showing cyt b_{150} , heme $b_{\rm H}$ and heme $b_{\rm L}$; the dashed line shows the curve expected for an *n*=1 component, with amplitude for degree of reduction plotted as $\Delta A = 0.055 \frac{1}{10^{(E_{\rm p}-E_{\rm m})/59}+1}$. (Bottom, right) spectra derived by fitting titrations at each wavelength using fixed values for $E_{\rm m}$, and *n*, for each component; (top, right) the pH dependence of $E_{\rm m}$ values for individual components, assuming single species for each component.



Fig. 7. Reanalysis of the thermodynamic behavior assuming the coulombic effects are in play. Titration data for heme *b*_H in the absence or presence of antimycin were assumed to reflect two components split by coulombic interaction.

complement of redox centers, except for that heme. The complex retained a fully functional Qo-site reaction, but turnover was restricted by the availability of heme $b_{\rm L}$ as the only low-potential chain acceptor [146]. Potentiometric titration of the hemes showed that the $E_{m,7}$ values for heme b_L were changed from ~-90 mV in wild-type to -8 mV (in H111N) or -14 mV (in H212D), an increase of ~80 mV. We suggested that this change could be explained by loss of a coulombic interaction on the loss of the negative charge of heme $b_{\rm H}$, which, with $E_{\rm m} \sim 50$ mV, would be reduced over the range in which heme $b_{\rm L}$ titrates. If this speculation is accepted, the value and the distances shown by structures would allow calibration of other coulombic interactions in the dimeric complex, and construction of the curves like those shown in Fig. 5. From this, it is evident that substantial coulombic effects might be expected both within the monomer, and across the dimer interface. Such interactions have been extensively investigated in other systems. In order to explore these possibilities, we have re-evaluated the thermodynamic properties of the b-hemes by analysis assuming such effects.

4.1. Thermodynamic properties of the b-hemes

Titrations were performed using full spectral analysis over the α - and β -band range, long equilibration times between measurements, and relatively high concentrations of mediators. Typical curves are shown in Figs. 6 (wild-type) and 8 (mutant N221I) [147]. Space does not

permit a full discussion, but several interesting features emerge from this new analysis.

- i) The pK at 7.8 previously observed is not seen [91,92]. It seems likely that in previous analysis, titration curves for heme $b_{\rm H}$ were convoluted with cyt b_{150} and the pH dependence of its amplitude (Fig. 6). Two brief reports suggest that the same is true in *Rb. capsulatus* [7,66].
- ii) A second new observation is the apparent independence of pH for the $E_{\rm m}$ values for heme $b_{\rm H}$ in the presence of antimycin.
- iii) Finally, in fitting the curves, especially those for $b_{\rm H}$ in the presence of antimycin, and for $b_{\rm L}$, we found that *n*-values less than 1 often provided better fits (Fig. 6, top left). When n=1 was imposed, analysis of the curves into two components was required for a good fit (Fig. 7).

Because small changes in the extremes of the fitted curves can give quite large differences in fitted values, the data presently available are not accurate enough in themselves to provide unambiguous values. Furthermore, analysis was complicated by the presence of additional components from other redox complexes, most notably a heme with a spectrum similar to heme $b_{\rm H}$ titrating in the range of the cyt $b_{\rm 150}$ component, which was not observed in titrations of the isolated complex, but which contributed 10–15% of the measured change in chromatophores. With these necessary caveats, when an analysis of

Table 4

Kinetic characteristics of strains with Asn-221 of cytochrome b modified

Strain	Photosynthetic growth	Reduction of $b_{\rm H}^{\rm a}$	Fraction of $b_{\rm H}$ reduced ^b	$E_{\rm m_{17}}$, pH dependence ^d			Re-oxidation $(t_1 c_2/ms)^e$
				Cyt b ₁₅₀	$b_{\rm H}$, no inhibitor	With AA	(*1/2/110)
Wild-type	+++	455	0.05	148, -59	32, -59	46, 0	<1.6
N221D	+++	(as wt)	0.8	150, -59	35, -54	-47, -59	15
N221H	+++	520	0.68	150, -59	35, -59	13, -50	12
N221P	+++	442	0.167	154, -59	38, -59	-3, -59	2.0
N221T	+++	(as wt)	0.37	140, -59	29, -50	13, 59	8.0
N221I	+++	(as wt)	0.75	104, -59	36, -59	-95, -59	14

^aRate of reduction of heme $b_{\rm H}$ measured following a 5 µs flash at $E_{\rm h} \sim 120$ mV in the presence of antimycin (units – mol cyt $b_{\rm H}$ /mol bc_1 complex/s).

^bFraction of heme $b_{\rm H}$ reduced at the maximal amplitude (at ~3–5 ms) of the kinetic trace in the absence of antimycin, compared to the fraction reduced in the presence of antimycin. ^cRate of re-oxidation of heme $b_{\rm H}$ in the absence of antimycin, measured by subtraction of the traces in the presence and absence of antimycin (units – mol cyt $b_{\rm H}$ /mol bc_1 complex/s). The rate in the wild-type reflects the rate-limiting reduction of the low-potential chain through the Q_o -site reaction. Since v = k(occupancy), and the fraction of heme $b_{\rm H}$ in the reduced form (column 4) is <<1, the rate constant must be much greater than that for the limiting step.

 ${}^{d}E_{m,7}$ in mV, pH dependence of E_{m} in mV/pH unit, assuming a single component for each heme.

^eHalf-times for re-oxidation of heme b_H measured from kinetic traces.

heme $b_{\rm H}$ was performed, assuming that each component of the bc_1 complex was split, some additional features became apparent (Fig. 7). The titration was resolved into two components in the presence or absence of antimycin, each showing the same pH dependence. The lack of pH dependence for heme $b_{\rm H}$ in the presence of antimycin disappeared, and was seen to reflect a pH dependence of the amplitude of a higherpotential component. The higher-potential component in the presence of antimycin had a lower $E_{\rm m}$ than the cyt b_{150} component, and so was merged with the low $E_{\rm m}$ component to give what appeared to be a single component. We hesitate to put too much emphasis on this reanalysis, but it suggests two populations with the same spectrum titrating with different $E_{\rm m}$ values, which is the behavior expected from two hemes in identical environments interacting through coulombic repulsion. Clearly, a more detailed analysis with full electrostatic calculations (cf. [69]) is needed, since the local fields and dielectric properties are not well represented by mean values assumed in Fig. 5.

4.2. Asn-221 mutants

Similar experiments with strains N221T, H, I, S, P and D have shown dramatic effects on the thermodynamic parameters measured in the presence of antimycin. Effects of mutation are also reflected in major changes in kinetics. These experiments are still in progress, but example data have been reported elsewhere [90,147], and are summarized in Table 4.

Several interesting effects can be noted in these data. In all Asn-221 mutant strains, the redox properties of both *b*-hemes (and the cyt b_{150} component) were relatively unaffected in the absence of antimycin, but $E_{\rm m}$ values for heme $b_{\rm H}$ were shifted to lower values in the presence of antimycin, quite dramatically in some strains. However, in the presence of antimycin, the electron transfer rates to the $b_{\rm H}$ heme were not markedly changed, indicating that turnover of the Q₀-site was unaffected. All mutations showed some inhibition of electron transfer from heme $b_{\rm H}$ to the acceptor at the Q_i-site in the absence of antimycin. Inhibition was apparent both at $E_h \sim 200$ mV, where the acceptor would be Q, and 100 mV, where SQ is likely the acceptor in WT. Titrations of SQ have shown similar values to wild-type for amplitude and peak of the bell-shaped titration curve in strains so far tested (N221P and S), so we have no reason to believe that the SO stability is markedly changed. For several strains, rates were substantially slower than wild-type. Surprisingly, N221P, in which H-bonding potential is lost, showed the least kinetic effect.

In the presence of antimycin, all strains showed changes in thermodynamic properties of heme $b_{\rm H}$ (but not $b_{\rm I}$). When analyzed according to conventional single component titrations, all mutants showed a dependence of ~-59 mV/pH. For several strains, the changes in $E_{\rm m}$ in the presence of antimycin were quite dramatic, as seen in N221I (Fig. 8). This can be seen most easily through the difference spectra in Fig. 8C. In the absence of antimycin, the difference ΔA for the redox cut between $E_{h.8.7}$ values -380 and -40 mV shows a spectrum of heme b_L similar to that over the same range in wild-type. On addition of antimycin, an additional component appears in this difference spectrum, shown as the antimycin induced change, which has the spectrum of heme $b_{\rm H}$, but with an $E_{\rm m}$ shifted to the same range as heme $b_{\rm L}$. As with wild-type, the titration data in the mutant strains were not well fit by single components of n = 1 (Fig. 8B), and this was also obvious in titrations of heme $b_{\rm L}$ (not shown). Because in some mutants, the two hemes showed similar $E_{\rm m}$ values, unambiguous resolution of the different components was difficult. The effects of the lowered E_m were apparent in kinetic data, so that in the presence of antimycin at ambient potentials at which the quinone pool was oxidized enough to sustain turnover ($E_{h,7}$ >40 mV), and where both hemes were expected to be substantially oxidized, reduction of components with both spectra was observed on flash activation. This is in contrast to wild-type, where reduction of heme $b_{\rm I}$ was observed only when heme $b_{\rm H}$ had been prereduced before the flash [109]. The interpretation of these results is



Fig. 8. Titration of heme $b_{\rm H}$ in chromatophores from N2211 in the presence and absence of antimycin. A (top), titration curves showing shift in $E_{\rm m}$ on addition of antimycin; B (middle), titration with antimycin fit by 2-component, n=1; C (bottom), absorbance spectra showing the difference (-380)–(-40) mV from pH 8.7 titration. The difference with and without antimycin shows the contribution of heme $b_{\rm H}$ with its low potential in the presence of antimycin.

again ambiguous, but we favor a picture in which coulombic interactions can occur both within the monomer, and across the dimer interface, as shown schematically in Fig. 9.

It will be obvious that detailed characterization of these strains, especially with respect to the SQ properties, will provide parameters suitable for testing the hypothesis proposed for cyt b_{150} formation. The dramatic effects of antimycin in the heme $b_{\rm H}$ properties will also help in understanding how binding can modify these parameters. A comprehensive review of structural aspects of antimycin binding by Huang et al.[86], the electrostatic calculations for the yeast complex [69], and the recent higher resolution structures of the *Rb. sphaeroides* complex [87] have provided a firm basis for discussion. Since the N221P strain showed minimal effects on function in the absence of

antimycin, it is clear that the H-bonding possibilities of the asparagine in wild-type are not essential to function [90]. Despite the minimal effect of mutation on function, on addition of antimycin this strain showed a marked shift in E_m for heme b_H. The pattern of this effect among the neutral residue changes tested was for a larger shift in E_m with increasing hydrophobicity. The N221D mutant falls outside this pattern, but this may reflect its acidic nature. There is no obvious feature of these characteristics that allows us at present to suggest a mechanistic explanation. Examination of the protein environment with respect to heme b_H shows a highly polar region, likely related to



Fig. 9. Scheme to show some of the coulombic interactions suggested. The dimer is represented by four Q-sites (circles) with antimycin blocking Q_i (blue cross). The hemes are shown as horizontal bars, either empty, or occupied by an electron (red dot). Top, in wild-type, heme b_H has $E_{m,7} \sim 50$ mV, heme b_L has in intrinsic $E_{m,7} \sim -30$ mV, but titrates with $E_m \sim -100$ mV because of coulombic interaction with heme b_H . Bottom, in N2211, both hemes have the same intrinsic $E_{m,7} \sim -30$. The first electron can go to either, and change the E_m of the other. For clarity only the coulombic interactions within a monomer are shown However, the titrations suggest that additional interactions across the dimer interface may also occur.

the need for equilibration of the heme with external pH. If such equilibration were through protonation of the D-propionate of heme $b_{\rm H}$ on redox change, this volume would be expected to support proton equilibration through H⁺ conducting pathways. We have noted previously the extensive H-bonding in this volume, and the presence of a bound water that forms a H-bonding link between the propionate and *Ser-203* in the bovine structure. Since there is considerable variability in conformation within this volume in the six different monomers of the *Rb. sphaeroides* bc_1 complex that are included in PDB file 2qjy, it seems likely that this part of the protein is structurally plastic. It may be involved in equilibration of the heme propionate with the N-phase pH. However, this propionate was not included in the set of residues shown to undergo protonation change in response to reduction in electrostatic studies [69].

In summary, it seems likely that coulombic interactions are "felt" between all centers in the dimeric structure that carry charge. The mechanistic consequences for interactions within a monomer are subsumed under the monomeric Q-cycle, since the E_m values derived from redox titration are those that reflect the interactions. As noted elsewhere, coulombic interactions within the Q_o-site provide a possible mechanism for prevention of bypass reactions [5]. Interactions across the dimer interface have not been analyzed in detail. However, it seems reasonable to look to an interaction between Q⁻⁻ species in the Q_i-site as a possible explanation for the otherwise anomalous observation that the maximal SQ population observed is a 1/dimer.

5. Conclusions

The modified Q-cycle continues to provide the most economical model for the mechanism of the bc_1 complex. The evidence for intermonomer electron traffic does not seem compelling, and our own experiments suggest that no such reactions occur on the time scale expected. Coulombic effects are expected, and can be demonstrated for the intra-monomer case. Analysis of titration curves suggests that coulombic interactions also occur across the dimer interface, as would be expected from simple physicochemical considerations.

Acknowledgments

We gratefully acknowledge support for the work reported here from grants NIH RO1 GM35438 (to ARC) and NIH RO1 GM62954 (to SAD).

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