Mitochondrial Disease-related Mutation G167P in Cytochrome b of Rhodobacter capsulatus Cytochrome bc1 (\$151P in Human) Affects the Equilibrium Distribution of [2Fe-2S] Cluster and Generation of Superoxide*

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Background: Mutation S151P was found in patients with exercise intolerance.

Results: Bacterial analogous substitution (G167P) influences movement of the iron-sulfur protein head domain (ISP-HD), increasing ROS production.

Conclusion: This correlation corroborates the recently proposed "semireverse" electron transfer mechanism of ROS

Significance: The molecular effect identified for S151P may be valid for several other human mutations that affect motion of ISP-HD.

Cytochrome bc_1 is one of the key enzymes of many bioenergetic systems. Its operation involves a large scale movement of a head domain of iron-sulfur protein (ISP-HD), which functionally connects the catalytic quinol oxidation Qo site in cytochrome b with cytochrome c_1 . The Q_0 site under certain conditions can generate reactive oxygen species in the reaction scheme depending on the actual position of ISP-HD in respect to the Qo site. Here, using a bacterial system, we show that mutation G167P in cytochrome b shifts the equilibrium distribution of ISP-HD toward positions remote from the Q_o site. This renders cytochrome bc_1 non-functional in vivo. This effect is remediated by addition of alanine insertions (1Ala and 2Ala) in the neck region of the ISP subunit. These insertions, which on their own shift the equilibrium distribution of ISP-HD in the opposite direction (i.e. toward the Qo site), also act in this manner in the presence of G167P. Changes in the equilibrium distribution of ISP-HD in G167P lead to an increased propensity of cytochrome bc_1 to generate superoxide, which becomes evident when the concentration of quinone increases. This result corroborates the recently proposed model in which "semireverse" electron transfer back to the Qo site, occurring when ISP-HD is remote from the site, favors reactive oxygen species production. G167P suggests possible molecular effects of S151P (corresponding in sequence to G167P) identified as a mitochondrial disease-related mutation in human cytochrome b. These effects may be valid for other human mutations that change the equilibrium distribution of ISP-HD in a manner similar to G167P.

Cytochrome bc_1 is a key component of the electron transport chain (for a recent review, see Ref. 1). It catalyzes the reaction of reduction of cytochrome c by quinol. Its action is related to building a proton-motive force, which is utilized to produce ATP. As a part of the mitochondrial electron transport chain, cytochrome bc_1 (complex III) plays a crucial role in oxidative phosphorylation (2).

In the catalytic Q cycle of cytochrome bc_1 (3, 4), energy connected with quinol to cytochrome *c* electron flow is utilized for proton translocation. In this case, electrons derived from the quinol oxidation are split at the Q_o catalytic site into two different cofactor chains: high potential c-chain (which includes Rieske cluster, heme c_1 , and heme c embedded in iron-sulfur protein (ISP),³ cytochrome c_1 , and diffusible substrate cytochrome c, respectively) and low potential b-chain (which includes heme b_L , heme b_H , and the quinone reduction Q_i site embedded in cytochrome b subunit) (Fig. 1A). Electrons transferred via the b-chain reduce quinone in the Qi site in two sequential steps (5). Transfer of electrons via the c-chain is made possible by movement of the head domain of iron-sulfur protein (ISP-HD), an important mechanistic element of the cytochrome bc_1 . ISP-HD moves between a site on the cytochrome b interface close to the Q_0 site (Q_0 position) and a site on the cytochrome c_1 interface (c_1 position) (6–11). Unstable semiquinone is frequently discussed as an intermediate of the Q_o site reaction (12–16). Recent experiments have trapped

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³ The abbreviations used are: ISP, iron-sulfur protein; ISP-HD, head domain of iron-sulfur protein; WT, wild type; O₂, superoxide ion; SOD, superoxide dismutase; CuZn-SOD, copper- and zinc-containing superoxide dismutase; Bicine, N,N-bis(2-hydroxyethyl)glycine; CW, continuous wave; E_{m8} , midpoint redox potential at pH 8.0; ROS, reactive oxygen species.

semiquinone in this site in the state of a free radical (17-19) or as a semiquinone coupled to the reduced Rieske cluster (19).

However, under certain conditions, part of the energy released from oxidation of quinol can be dissipated through side reactions that result in partial or total loss of proton translocation function of cytochrome bc_1 . These reactions reduce the efficiency of separation of electrons in the Q_o site into two cofactor chains. One of the possible side reactions is an electron transfer from semiquinone generated in the Q_o site to oxygen to produce superoxide (15, 20, 21). This reaction causes a decrease in the yield of proton translocation and results in formation of free radicals that could cause damage to certain macromolecules including lipids and proteins.

Under some specific conditions, when the electron flow through cofactor chains is impeded, the side reactions in which superoxide is generated may be enhanced. For example, the presence of antimycin blocks the electron flow through the Q_i site (20–23) and favors reverse electron transfer from heme b_L to quinone with formation of semiquinone in the Q_o site (24, 25). The formed semiquinone can react with ISP if ISP-HD is at the Q_o position, or it can react with oxygen if ISP-HD is not present in the Q_o site. Therefore states of the Q_o site with ISP-HD not present at this site favor superoxide production (25, 26).

Among all subunits of complex III, only cytochrome b is encoded by mtDNA. Thus the probability of the occurrence of mutations in this subunit is higher compared with other subunits of the complex encoded by nDNA. In principle, such mutations may have various effects on function including adaptive effects proposed for mutations in humans: T15204C (I153T), T14798C (F18L), and G15257A (D171N) found in haplogroup C, J1, and J2, respectively (27). Nevertheless, most of the mutations identified so far in human cytochrome b appear to be linked with diseases such as exercise intolerance, myopathy, and cardiomyopathy (28-31). They all are somehow related to reduced efficiency of energy conversion, which could be caused by altered operation of mutated complex III. However, studying the molecular effects of these mutations in humans is often difficult, especially in the context of the occurrence of heteroplasmy and limited amount of protein that can be obtained from human cells. Those types of limitations can be overcome when using bacterial (32, 33) or yeast (30, 34) systems, which thus provide a good model to study human mitochondrial disease-related mutations at the molecular level. Conversely, the identified effects of certain human mutations can be a valuable source of general information on mechanisms of catalytic and side reactions.

This approach was undertaken in the present study. We chose mutation T15197C (in the mtDNA sequence) identified in patients with exercise intolerance (35). This mutation changes serine 151 to proline in the vicinity of the Q_o site and was found to result in expression of complex III with slightly reduced amounts of cytochrome b and cytochrome c_1 (35). In a yeast model, the analogous mutation (S152P) resulted in expression of complex III with significantly reduced amounts of ISP subunit (29). We show that this mutation introduced at an analogous position to the cytochrome b subunit (G167P) of bacterial cytochrome bc_1 influences the average position of

ISP-HD with respect to other subunits in such a way that it stays more remote from the Q_o site than in the native enzyme. At the same time, no changes in the amounts of subunits in the catalytic core were observed. This provided new insight into the possible molecular basis of the human disease associated with the presence of T15197C. It also provided the first (to our knowledge) mutational variant of cytochrome bc_1 in which a shift of ISP out of the Q_o site was identified spectroscopically. This allowed us to extend the array of conditions that tested changes in the levels of superoxide production by cytochrome bc_1 toward a better understanding of the mechanism of this reaction.

Experimental Procedures

Preparation of Mutants-To construct mutations in cytochrome bc_1 , a genetic system originally developed by Dr. F. Daldal (University of Pennsylvania, Philadelphia, PA) (36) was used. G167P mutation (underlined) was introduced in the gene coding for cytochrome b (petB) using the QuikChange sitedirected mutagenesis system (Stratagene) and the following PCR primers: G167P_FWD (5'-C ACC GGC CTG TTT CCG GCG ATC CCG GGC ATC G-3') and G167P_REV (5'-GCC CGG GAT CGC CGG AAA CAG GCC GGT GAT CAC G-3'). To construct single mutant G167P, pPET1 plasmid containing wild type (WT) petABC operon was used as a template DNA. To obtain G167P/1Ala and G167P/2Ala double mutants, the template DNA contained appropriate GCG insertions (resulting in 1Ala or 2Ala) in the petA gene coding for ISP (10). The correct sequence of engineered constructs was verified by sequencing entire *petA* and *petB* genes. The BstXI-XmaI fragments of the operon containing the desired mutations and no other mutations were exchanged with WT counterpart of pMTS1 plasmid. Expression vectors were introduced into MT-RBC1 Rhodobacter capsulatus strain (devoid of petABC operon) using triparental crossing (36). The presence of introduced mutations was confirmed by sequencing the petA and petB genes on a plasmid isolated from the mutated R. capsulatus strains.

 $R.\ capsulatus$ bacteria were grown under semiaerobic or photoheterotrophic conditions as described previously (37). To test for the occurrence of reversion mutations, $100\ \mu l$ of 2 ml of overnight liquid culture of the mutant strains were spread on mineral-peptone-yeast extract (MPYE) plates and kept in selective photosynthetic cultures for 10 days. Single colonies that acquired the Ps⁺ phenotype (photosynthetic competence) were isolated, and reversion mutations were identified by sequencing the entire petABC operon.

Isolation and Purification of Cytochrome bc_1 Complexes—Chromatophores from R. capsulatus cells grown under semiaerobic conditions were obtained using the procedure described previously (38). Cytochrome bc_1 complexes were isolated from detergent-solubilized chromatophores using ion exchange chromatography (DEAE-BioGel A) as described (38).

Steady-state Kinetics Measurements—Steady-state enzymatic activity of isolated cytochrome bc_1 complexes was determined spectroscopically by the 2,3-dimethoxy-5-methyl-6-decyl-1,4-benzohydroquinone-dependent reduction of cytochrome c (bovine heart cytochrome c from Sigma-Al-



drich) as described previously (36). All enzymatic assays were performed in 50 mm Tris buffer (pH 8) containing 0.01% n-dodecyl β -D-maltoside and 100 mm NaCl. The final concentrations of substrates were as described in Table 1 and the legend to Fig. 8. Concentrations of cytochrome bc_1 were in the range of 10-100 nm depending on activity of the mutant. Turnover rates were calculated from the initial linear parts of the curves. The level of superoxide production was expressed as percent difference of enzymatic activity in the presence and absence of 100 units/ml CuZn-SOD (20, 25, 26). Student's t test for independent samples was used for statistical analysis. Only p values lower than 0.05 were considered statistically significant.

Superoxide radical production was also measured via hydrogen peroxide formation using the Amplex Red horseradish peroxidase method (Amplex Red Hydrogen Peroxide/Peroxidase Assay Kit, Life Technologies). Assays were performed in 50 mm Tris buffer (pH 8) containing 0.01% n-dodecyl β -D-maltoside and 100 mm NaCl. The final concentrations of substrates were as described in the legend to Fig. 9. The reaction mixture also contained 50 µM Amplex Red reagent, 0.1 unit/ml HRP, and 300 units/ml CuZn-SOD. CuZn-SOD was added in excess to convert superoxide into hydrogen peroxide, which in the presence of horseradish peroxidase reacts with Amplex Red reagent to produce the red fluorescent oxidation product resorufin. Resorufin has a broad absorption peak (between 500 and 590 nm) with maximum at 571 nm. The increase in resorufin absorbance was followed at the isosbestic point of cytochrome c at 540 nm. The levels of H₂O₂ were determined from the absorbance at 540 nm measured at the time point when all substrates were used up (after reduction of 10 μ M cytochrome c).

Light-induced Electron Transfer Measurements—Double wavelength time-resolved spectrophotometry (39) was used to obtain the transient kinetics of heme b reduction at 560-570 nm. All measurements were performed using bacterial chromatophores suspended in MOPS buffer (pH7) containing 1 mm EDTA and 100 mm KCl following the procedure described previously (39, 40). The samples were poised at an ambient potential of 100 mV in the presence of $3.5~\mu{\rm M}$ valinomycin and redox mediators as described (39). Rates of flash-induced heme b reduction were determined by fitting transient kinetics to a single exponential equation.

[2Fe-2S] Cluster Relaxation Measured by Pulse EPR—The temperature dependence of phase relaxation rate of [2Fe-2S] cluster was measured in isolated complexes using pulse EPR spectroscopy. The measurements were carried out on a Bruker Elexsys-E580 spectrometer at Q-band (33.5 GHz). The electron spin echo decay of each sample was recorded in temperature range from 12 to 24 K in the same manner as described previously (41). The relaxation rates were determined from fitting a stretched exponential function to the measured electron spin echo curves. The samples were prepared in 50 mm Bicine buffer (pH 8) and 100 mm NaCl under reducing conditions (1 mm sodium ascorbate) in the presence of 20% glycerol as described previously (41). The measured relaxation rates concern intracomplex interactions and are highly reproducible, falling within a standard error typical of fitting procedure (1-2%), irrespective of protein isolation.

The EPR Potentiometric Titration of [2Fe-2S] Cluster— Potentiometric titrations of [2Fe-2S] cluster in chromatophore membranes were conducted as described (40, 42). Measurements were performed in 50 mm Bicine buffer (pH 8) containing 100 mm KCl, 20% glycerol, and mediators 2,3,5, 6-tetramethyl-1,4-phenylenediamine, 1,2-naphthoquinone-4sulfonate, 1,2-naphthoquinone, 2,3,5,6-tetrachlorohyroquinone, N-ethyldibenzopyrazine ethyl sulfate salt, and N-methylphenazonium methyl sulfate, each at a concentration of 100 μM. All CW EPR spectra were recorded at 20 K using the following parameters: power, 1.9 mW; frequency, 9.39 GHz; modulation amplitude, 10 G. To obtain the value of midpoint redox potential at pH 8.0 (E_{m8}), the amplitude of CW EPR spectra of the reduced [2Fe-2S] cluster was plotted as a function of ambient redox potential (E_h) and fitted with the Nernst equation assuming a one-electron couple.

Results

Spectral and Kinetic Properties of G167P in Cytochrome b— As shown in Fig. 1C, Gly-167 in R. capsulatus cytochrome b is located at the end of helix cd1 close to the twist loop that separates this helix from helix cd2. This region is at the entry of the Q_o quinol binding pocket and comes in close contact with ISP-HD when it approaches cytochrome b. In mitochondrial complex III, the corresponding position is occupied by Ser (Ser-151). Introducing Pro at this position (G167P) results in expression of cytochrome bc_1 in bc_2 capsulatus cells that contains all catalytic subunits (cytochrome bc_1 , cytochrome bc_2 , and ISP) as visualized on SDS gels (Fig. 2 bc_2).

However, G167P mutant is not functional *in vivo* as indicated by the incapability of cells to sustain cytochrome bc_1 -dependent photosynthetic growth (Fig. 3 and Table 1). This correlates with low enzymatic activity of the mutant (turnover rate of 14.7 versus 140 s⁻¹ in WT) and severe impediments in the operation of the Q_o site identified by light-induced kinetic measurements. As shown in Fig. 4 (A and B, red trace), the Q_0 site-mediated reduction of hemes b in the presence of antimycin is greatly inhibited in G167P (Table 1). In the absence of inhibitors, the reduction of hemes b through the Q_0 site and reoxidation through the Q_i site takes place (Fig. 4, A and B, black trace) but because of the much slower rate of heme b reduction is not kinetically resolved as in WT and only visualized as the difference between the black and red traces in Fig. 4B. Overall, the enzymatic assays and light-induced measurements indicate severe, but not complete, inhibition of the Q_o site in G167P.

The optical spectra of hemes b and c in the G167P mutant do not differ from the spectra of native complex (Fig. 2B). They exhibit an ascorbate-reducible peak at 553 nm (reflecting high potential heme c_1) and dithionite-reducible peaks at 553 and 560 nm (the latter one reflecting the low potential hemes b_L and b_H). Conversely, the CW EPR spectra of the Rieske cluster in G167P clearly differed from that of the cluster in WT cytochrome bc_1 (Fig. 2C): the g_x transition in the mutant is much broadened and of a smaller amplitude as compared with the sharp and characteristic $g_x = 1.805$ present in WT. In addition, the g_x in G167P appears to be composed of more than one component (see also the spectrum of G167P in Fig. 7). This change in the shape of the spectrum provides the first indica-

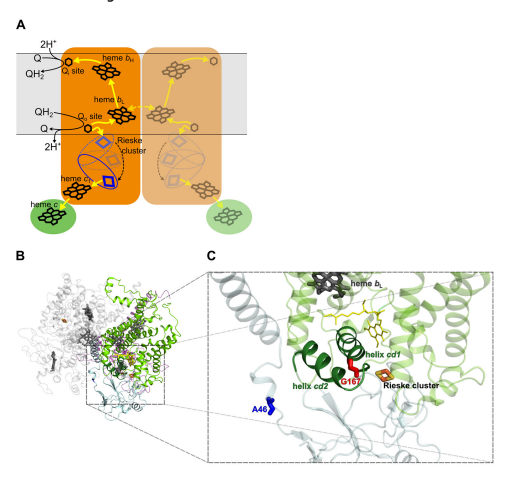


FIGURE 1. A, a simplified scheme of reactions occurring in dimeric cytochrome bc_1 . Cytochrome bc_1 catalyzes reduction of cytochrome c (green ellipse) by quinol. The Q cycle reactions are highlighted for one monomer (left, orange shape). Yellow arrows designate the electron transfer pathway. For simplicity, only forward reactions are shown. The yellow dotted arrow represents intermonomer electron transfer between the b_L hemes (57, 58). A domain harboring Rieske cluster (blue) moves (black dotted arrow) between Q_0 position and c_1 position. The membrane is shown as a light gray area. B, crystal structure of dimeric cytochrome bc_1 from B. capsulatus (Protein Data Bank code 1ZRT (59)). Subunits of one monomer are colored as follows: cytochrome b, light green; ISP, gray; cytochrome c_1 , transparent blue. Subunits of the second monomer are in light gray. C, Close up view of the part of cytochrome b and ISP showing structural details of the Q_0 site. Gly-167 (red sticks) in cytochrome b (light green) is located in the end of helix cd1 (dark green). Ala-46 (blue sticks) located in the neck region of ISP (gray) indicates the position where one (1Ala) or two (2Ala) alanine residues were inserted. Dark gray and light orange sticks indicate heme b_L and Rieske cluster, respectively. Yellow lines represent the stigmatellin molecule.

tion that the interaction between the Rieske cluster and quinone in the $Q_{\rm o}$ site is altered due to changes in the occupancy of the site with quinone and/or changes in the position of ISP-HD caused by structural constraints affecting the motion of this domain. Those types of changes would provide an explanation for the enzymatic and kinetic impediments observed in G167P. The g_x transition remains sensitive to the addition of $Q_{\rm o}$ site-specific inhibitors stigmatellin and myxothiazol (Fig. 2C). The shape of g_x in the presence of these inhibitors in G167P is similar to that of native complex: in both cases, stigmatellin induces sharp g_x at 1.783, whereas myxothiazol largely broadens this transition, shifting it to a value of $g_x=1.763$.

Phase Relaxation of Rieske Cluster in G167P—To get further insights into the molecular effects of G167P, we performed analysis of the temperature dependence of the phase relaxation rate of the Rieske cluster. Our previous studies showed that changes in phase relaxation rate reflect changes in the position of ISP-HD (41). More specifically, at the macroscopic level of the protein solution, they reflect changes in the average equilibrium position of ISP-HD with respect to other subunits of the

complex. This approach benefits from the observation that oxidized heme $b_{\rm L}$ enhances the relaxation in a distance-dependent manner: the closer the [2Fe-2S] cluster to heme $b_{\rm L}$, the stronger the enhancement. This means that the movement of ISP-HD out of the ${\rm Q_o}$ site resulting in an increase of distance between heme $b_{\rm L}$ and Rieske cluster weakens the enhancement (41). A comparison of the temperature dependence profiles of the phase relaxation rate of the Rieske cluster shown in Fig. 5A reveals that the enhancement in G167P is weaker compared with WT. This indicates that in G167P the average equilibrium position of ISP-HD is set at a larger distance from heme $b_{\rm L}$ (and the ${\rm Q_o}$ site) than in WT (Fig. 6, *right versus middle*). In other words, G167P causes a shift in the equilibrium position of ISP-HD toward positions more remote from the ${\rm Q_o}$ site.

Reversions—When incubated under photosynthetic conditions, a photosynthetically inactive strain carrying G167P reverts spontaneously to photosynthetically competent cells (see Fig. 3). The DNA analysis of the revertant cells revealed that they are same site revertants and replace Pro at 167 with Ser or Gln (of eight clones analyzed, seven introduced Ser (gen-

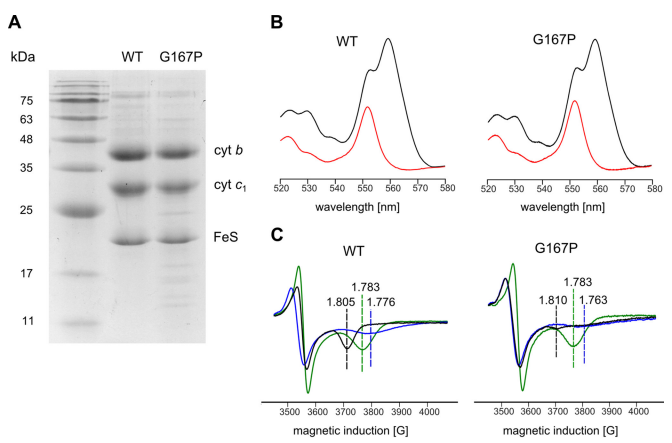


FIGURE 2. **Spectral properties and subunit composition of WT and G167P mutant.** A, SDS-PAGE analysis of isolated complexes. B, optical difference spectra of purified cytochrome (C) C0 complexes. C1 complexes. C3 complexes. C4 and C7 complexes. C8 correspond to dithionite minus ferricyanide and ascorbate minus ferricyanide spectra, respectively. C7 CN band CW EPR spectra of the [2Fe-2S] cluster measured in chromatophores suspended in 50 mm MOPS (C1 and 100 mm KCl. Samples were reduced with ascorbate in the absence of any inhibitor (C8 and in the presence of myxothiazol (C9 or stigmatellin (C9 are traces). C9 Dotted lines indicate the position of the C9 transition. C9 gauss.

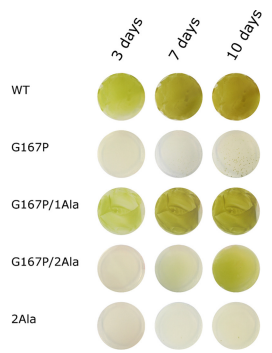


FIGURE 3. **Photosynthetic growth of various** *R. capsulatus* **strains.** Single *green dots* visible on the G167P plate after 10 days represent revertants (Ps⁺). Panels show growth on plates observed after 3, 7, and 10 days.

erating mutant G167S) and one introduced Gln (generating mutant G167Q)). The enzymatic assays and kinetic data indicated that both G167S and G167Q regain much of their electron transfer activity compared with the original G167P. However, compared with WT, G167Q still exhibits about 2 times slower enzymatic turnover and rate of flash-induced heme *b* reduction, whereas G167S in both measurements approaches the WT level (Table 1). It is of note that position 167 is naturally occupied by Ser in several species including humans (corresponding position is Ser-151). High electron transfer activity of G167S is consistent with this observation.

Effects of Combination of G167P in Cytochrome b with 1Ala and 2Ala Insertions in the Neck Region of ISP Subunit—It was previously recognized that alanine insertion (1Ala or 2Ala) in the neck region connecting the ISP-HD with its hydrophobic anchor introduces steric constrains to the motion of ISP-HD (10). As a result, ISP-HD stays captured at the Q_o site for milliseconds in 1Ala mutant or seconds in 2Ala mutant, which is a much longer time than in WT (microseconds or less) (10). Therefore, in solution containing 1Ala or 2Ala mutants, there is an increase in the population of complexes having ISP-HD at the Q_o site (25, 41). In this context, a direction of the shift in the average position of ISP-HD caused by G167P can be considered opposite to the one caused by 1Ala or 2Ala mutants (Fig. 6). This prompted us to analyze the effect of a combination of G167P and alanine insertions in two double mutants, G167P/

TABLE 1Selected properties of *R. capsulatus* mutants

Strains	Phenotype ^a	Enzymatic activity ^b	Flash-induced heme b reduction ^c	E_{m8} of Rieske cluster	Reversions
		s^{-1}	s^{-1}	mV	
WT	Ps^+	140 ± 5	818	308	NA^d
G167P	Ps ⁻	14.7 ± 0.2	16.6	290	G167Q, G167S
G167P/1Ala	Ps ⁺	34 ± 2	131	293	NA
G167P/2Ala	Ps ^{slow}	21 ± 1	22.6	300	NA
1Ala	Ps ⁺	144 ± 7	101	334	NA
2Ala	Ps ⁻	4.2 ± 0.2	ND^e	390	ND
G167Q	Ps ⁺	70 ± 4	472	ND	NA
G167S	Ps ⁺	105 ± 5	805	ND	NA

^a Ps⁺ and Ps⁻ indicate photosynthetic competence and incompetence, respectively.

^e ND, not determined.

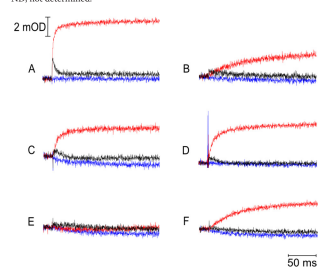


FIGURE 4. **Flash-activated heme** *b* **reduction.** The traces were recorded for WT (*A*), G167P (*B*), 1Ala (*C*), G167P/1Ala (*D*), 2Ala (*E*), and G167P/2Ala (*F*) at pH 7 and an ambient potential of 100 mV. Kinetic transients at 560 –570 nm were recorded without inhibitors (*black lines*) and in the presence of antimycin or myxothiazol (*red* or *blue lines*, respectively). *mOD*, milli-optical density units.

1Ala and G167P/2Ala. As shown in Fig. 3 and Table 1, both mutants, unlike G167P, can grow under photosynthetic conditions. Clearly, either 1Ala or 2Ala can suppress effects caused by G167P. However, the growth of G167P/1Ala is more vigorous than that of G167P/2Ala, indicating that the former combination of mutations yields cytochrome bc_1 that operates more efficiently. This difference finds its roots in the different phenotypic properties of the original Ala insertions (10): 1Ala is photosynthetically active, whereas 2Ala is not (Table 1). Thus, although in the case of G167P/1Ala an addition of the Ps⁺ mutation (1Ala) to the Ps⁻ mutation (G167P) rendered the double mutant Ps⁺, in the case of G167P/2Ala, a weak Ps⁺ phenotype was achieved by combining the two mutations that originally were Ps⁻ (Fig. 3 and Table 1).

The enzymatic activities and measured rates of light-induced electron transfer in the double mutants seem consistent with their phenotypic properties (Fig. 4 and Table 1). G167P/2Ala shows a slight increase in the turnover rate and rate of light-induced heme b reduction as compared with G167P. However, changes in G167P/2Ala are more dramatic in relation to the 2Ala mutant, which has a very low enzymatic activity (turnover of around 4 s $^{-1}$) and no signs of millisecond heme b reduction

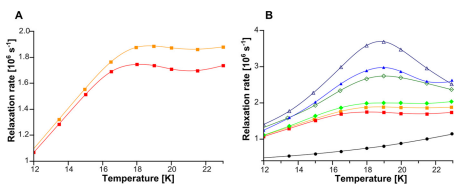
under the conditions described in Fig. 4 (G167P/2Ala displays a 5 times higher turnover rate and a clear millisecond reduction of heme b). G167P/1Ala shows a further increase in both the enzymatic activities and heme b reduction compared with G167P/2Ala (Table 1). However, the extent of the increase is clearly larger for heme *b* reduction than for enzymatic activity. The same trend is also observed when G167P/1Ala is compared with G167P: the rate of flash-induced heme b reduction increases about an order of magnitude, whereas the enzymatic activity increases only 2 times (Table 1). In fact, the rate of heme b reduction in G167P/1Ala reaches the level observed in 1Ala; however, the enzymatic activity of 1Ala is about 4 times higher compared with G167P/1Ala. This makes G167P/1Ala an interesting example of a combination of mutations exerting more severe inhibitory effects under the conditions of multiple turnover than under the conditions of flash-induced electron transfer.

As seen in the EPR spectra of Fig. 7, the heterogeneity of g_x transition originally observed in G167P is more pronounced in G167P/1Ala and G167P/2Ala. The g_x in EPR spectra of both double mutants allows for the distinction of two transitions (1.805 and 1.772). If $g_x = 1.805$ is taken into account, one can see an increase in the amplitude in the following order: G167P, G167P/1Ala, G167P/2Ala, WT. As the characteristic shape of $g_x = 1.805$ in WT is usually assigned as reflecting interaction of reduced [2Fe-2S] cluster with quinone bound in the Q_o site (43, 44), the increase in the amplitude of $g_x = 1.805$ in double mutants in comparison with G167P suggests that in those mutants the population of ISP-HD occupying the Q_o position is larger. This seems to follow the expectations based on the opposing effects of 1Ala or 2Ala and G167P. However, it is important to emphasize that the shape of the EPR spectrum in chromatophores frozen without glycerol may not necessarily reflect the distribution of positions of ISP-HD as best evidenced in 1Ala and 2Ala mutants, which both have EPR spectra similar in shape to WT but exert large effects on distribution of ISP-HD positions (Figs. 5B and 7) (25, 41, 45). Conversely, the increase in g_x in double mutants points toward the possibility that the change in the shape of EPR spectrum of G167P alone (Fig. 7) is caused by the remoteness of ISP-HD from the Q_o site rather than the absence of quinone bound at the Q_o site (provided that 2Ala or 1Ala does not change the affinity of Qo site for its substrate in G167P).

^b Enzymatic activity rates are expressed as μmol of cytochrome c reduced/μmol of cytochrome bc₁/s. Conditions were 50 mm Tris-HCl (pH 8), 100 mm NaCl, 20 μm quinol, and 20 μm oxidized cytochrome c from bovine heart. Errors represent S.D. of the mean of at least four measurements.

^c Flash-induced heme *b* reduction rates for pH 7 and ambient potential of 100 mV.

 $[^]d$ NA, not applicable.



 $FIGURE 5. \textbf{Temperature dependence of phase relaxation rate of \textbf{[2Fe-2S] cluster in native and mutated cytochrome \textbf{\textit{bc}}_{1}.A, G167P (\textit{red closed squares}) and \textbf{\textit{bc}}_{2}.A, G167P (\textit{red closed squares}) and \textbf{\textit{bc}}_{3}.A, G167P (\textit{red$ WT (orange closed squares). B, 2Ala (blue open triangles), G167P/2Ala (blue closed triangles), 1Ala (dark green open diamonds), G167P/1Ala (green closed diamonds), G167P (red closed squares), and WT (orange closed squares) (last two sets of data are replotted from A). Control sample (black closed circles) is WT in which there are no dipolar interactions between heme b_{\perp} and [2Fe-2S] cluster. All samples in A and B were reduced with ascorbate except for the control of WT that was reduced with dithionite.

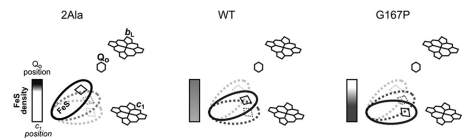


FIGURE 6. Scheme illustrating difference in equilibrium distribution of ISP-HD position in 2Ala and G167P in comparison with WT. In 2Ala mutant, ISP-HD stays captured at the Q_0 site for seconds, and its average position is shifted toward the Q_0 site in relation to WT. In the native cytochrome bc_1 , ISP-HD moves freely between Q_0 and c_1 positions. G167P has an effect opposite to 2Ala: the average position of ISP-HD is more remote from the Q_0 site than in WT. Horizontal rectangles depict the density of [2Fe-2S] cluster at the Q_o site: the darker and lighter shades denote higher and lower occupancy of the [2Fe-2S] cluster, respectively.

To get insights into the effects of G167P/1Ala and G167P/ 2Ala on the distribution of positions of ISP-HD, we compared the temperature dependence profiles of the phase relaxation rate of the Rieske cluster in those mutants, taking the profiles of 2Ala, 1Ala, and WT as a reference (Fig. 5B). The strongest enhancement observed in the 2Ala mutant reflects almost the entire population of ISP-HD at the $Q_{\rm o}$ site. The enhancement in G167P/2Ala is weaker, indicating a decrease in the population of ISP-HD at the Q_o site in this mutant. However, it stays above the level of enhancement seen in the 1Ala mutant, indicating that the population of ISP-HD at the Qo site in G167P/2Ala is larger than in 1Ala. The enhancement in G167P/1Ala is weaker than in 1Ala, indicating a further decrease in the population of ISP-HD at the Q_o site. This population is still larger than in WT (enhancement in G167P/1Ala is larger).

Overall, the decrease in the strength of the enhancement indicates a decrease in the population of ISP-HD at the Q_0 site and a shift in equilibrium position toward positions more remote from the Q_o site. It appears as if on one hand 1Ala and 2Ala diminished the "pushing" of the ISP-HD out of the Q_o site caused by G167P, and on the other hand, G167P diminished the pushing of the ISP-HD toward the Qo site caused by 1Ala and 2Ala. As a result, the average ISP-HD position in G167P/1Ala or G167P/2Ala became closer to the $Q_{\rm o}$ site than in WT (and G167P) but did not reach proximity to the Q_o site achieved in 1Ala or 2Ala, respectively. Considering all these effects, one can rank the mutations in the following order (with the first having the largest population of ISP-HD at the Q_o site): 2Ala > G167P/ 2Ala > 1Ala > G167P/1Ala > WT > G167P.

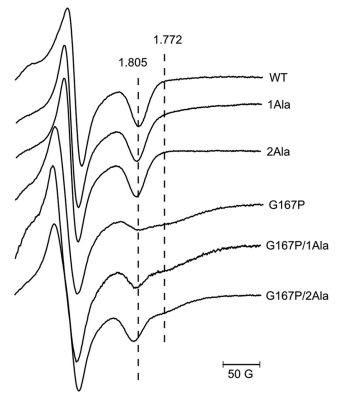


FIGURE 7. X-band CW EPR spectra of [2Fe-2S] cluster of WT and various mutants. All EPR spectra were measured in chromatophores suspended in 50 mm MOPS (pH 7) and 100 mm KCl and reduced with ascorbate. Dotted lines indicate the position of [2Fe-2S] g_x transitions.

Table 1 shows that values of midpoint redox potential (E_m) of the Rieske cluster in the mutants G167P, G167P/1Ala, and G167P/2Ala are similar to that of WT. This differentiates these mutants from the 1Ala and 2Ala mutants, which were previously shown to display an increase in E_m of Rieske cluster in respect to WT (in particular, 2Ala displays an E_m value about 100 mV higher than WT) (10). Given that the Ala mutants generally increase the population of ISP-HD at the Q_o site, their effect on E_m is consistent with the notion that the Rieske cluster in/close to the Q_o site has a higher E_m if compared with its E_m at other positions (46) (an increase in E_m is also observed in the presence of some inhibitors such as stigmatellin (47) that fix the ISP-HD at the Q_o site). In this context, the lack of an increase in E_{m8} of G167P, G167P/1Ala, and G167P/2Ala can be considered as additional support to the notion that a population of ISP-HD at the Q_o site is generally decreased in all G167P mutants (i.e. even when Ala insertions are present). We note, however, that the origin of the changes in E_m of the Rieske cluster is a complex issue as E_m not only depends on the position of ISP-HD but on several other factors including occupancy of the Q_o site with substrate/inhibitor (45, 48). Thus, a change in a value of E_m is not a simple translation to changes in ISP-HD position. A good example is provided by G167P/2Ala, which has a larger population of ISP-HD at the Q_0 site than 1Ala (Fig. 5B) but E_{m8} of Rieske cluster is not elevated (Table 1).

Effect of G167P on Generation of Superoxide at the Q_o Site of Cytochrome bc_I —It was previously recognized that superoxide is generated at the Q_o site in reactions involving a back electron transfer from heme b_L to quinone (24–26). This leads to the formation of semiquinone, which has the highest probability of reacting with oxygen when ISP-HD occupies a position remote from the Q_o site (25, 26). It follows that mutations shifting the average equilibrium position of ISP-HD out of the Q_o site should in principle enhance superoxide production by cytochrome bc_1 . G167P seemed a good candidate to test this assumption.

Typically, production of superoxide in native cytochrome bc_1 is observed when the inhibitor antimycin blocks the reoxidation of heme $b_{\rm H}$ (and $b_{\rm L}$) via the Q_i site (20, 26, 49). The measurements are performed in detergent solution with isolated enzyme exposed to an excess of substrate, cytochrome c and quinol. Our initial measurements indicated that under those types of conditions the G167P mutant generates even larger amounts of superoxide than the native cytochrome bc_1 (30 versus 17% for G167P and WT, respectively). This prompted us to verify whether this mutant can generate superoxide in the absence of any inhibitor. However, the measurements performed with an excess of quinol (typical conditions of enzymatic assays) revealed only a little superoxide generated by G167P (less than 5%). We thus tested a variety of conditions where both quinol and quinone were present in the reaction mixture in various proportions. The results are shown in Fig. 8. The amount of superoxide generation is described "quantitatively" as a number of μ mol of $O_2^{\overline{}}$ produced by 1 μ mol of the enzyme per second (Fig. 8B) or "relatively" as a percentage of superoxide generated per single turnover (Fig. 8C). Quantitatively, the superoxide generation shows a bell shape with the maximum at a quinol to quinone pool ratio of \sim 70 to 30%. Changes of this proportion in any direction lead to a decrease of the total amount of superoxide generated per second. Relatively, a linear increase of superoxide generation is observed upon an increase of quinone and decrease of quinol. The bell shape of O_2^- generation (Fig. 8*B*) is a consequence of a balance between two opposite effects associated with oxidation of the Q pool: O_2^- per single turnover increases with increasing concentration of quinone, while at the same time, the total enzymatic turnover rate decreases (Fig. 8*A*).

Under the conditions of maximum superoxide production per second observed for G167P, WT and mutants 2Ala and G167P/2Ala did not generate superoxide within experimental uncertainty (\pm S.E.) (Fig. 8*D*). The difference between mean values of enzymatic activities with and without SOD is shown for G167P (p value = 2×10^{-6} , statistically significant result) and G167P/2Ala (p value = 0.22, not statistically significant result). The results for WT and 2Ala are consistent with previous observations described (26).

To confirm enhanced production of superoxide in G167P mutant, we performed additional analysis using the Amplex Red horseradish peroxidase method (Fig. 9). We also observed that uninhibited G167P mutant produces a high level of superoxide, which was reflected in 10 times higher concentration of $\rm H_2O_2$ measured for this mutant with respect to WT, 2Ala, or G167P/2Ala. We note that the background level of $\rm H_2O_2$ observed for WT, 2Ala, and G167P/2Ala is consistent with the results reported for uninhibited complex III (24).

Discussion

Replacement of Gly-167 in cytochrome b with Pro is expected to result in structural distortions in the region encompassing helix cd1 and helix cd2, for example in a change in relative position of these two helices. As this region forms a part of the interaction site of cytochrome *b* with ISP-HD (7, 50, 51), such distortions may influence the process of binding/release of ISP-HD upon its interaction with cytochrome b. This will impact the motion of ISP-HD, which will manifest itself in the change in the equilibrium position of ISP-HD. Indeed, the relaxation enhancement measurements revealed that the average equilibrium position of ISP-HD in G167P is at a larger distance from heme $b_{\rm L}$ if compared with WT (Fig. 5A). This provides the first (to our knowledge) example of a mutation in which the shift of position of ISP-HD out of the Qo site was documented spectroscopically. So far, the documented and well characterized cases concerned mutations that shift the average position of ISP-HD in the opposite direction; i.e. the ISP-HD is arrested at the Qo site, which makes the average equilibrium position of ISP-HD at a closer distance from heme $b_{\rm L}$ (if compared with WT). The most prominent mutations that act in this way are insertions in the neck region of the ISP subunit (in particular 1Ala and 2Ala mutants, which arrest the ISP-HD for milliseconds or seconds, respectively) (10, 25, 41).

The effects of G167P on the distribution of ISP-HD caused a large decrease in the rates of electron transfer and enzymatic activity (Table 1), which rendered cytochrome bc_1 non-functional *in vivo* (the enzyme was not able to support growth of the cells under photosynthetic conditions) (Fig. 3). Interestingly, a simultaneous presence of the oppositely acting 1Ala or 2Ala

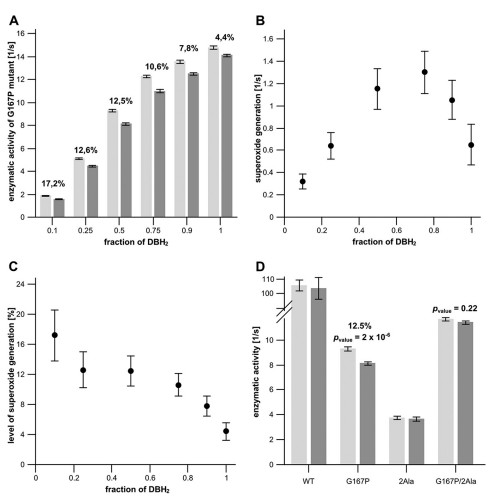


FIGURE 8. A, enzymatic activities of uninhibited G167P mutant under varying redox states of the quinone pool (quinol/quinone pool ratio) in the absence (*light gray bars*) and presence (*dark gray bars*) of SOD. Conditions were 50 mM Tris-HCl (pH 8), 100 mM NaCl, and 20 μ M oxidized cytochrome c. The total concentration of the quinone pool was 20 μ M. The amount of superoxide production is shown as a percentage *above* the *bars*. *Error bars* represent S.D. of the mean of 12 measurements. B, rates of superoxide generation by G167P calculated from the data in A. C, percentage of SOD-sensitive cytochrome c reduction in relation to cytochrome c reduction measured in the absence of SOD (based on the data of A). D, enzymatic activities of WT and mutants (G167P, 2Ala, and G167P/2Ala) measured in the absence (*light gray bars*) and presence (*dark gray bars*) of SOD. Conditions were 50 mM Tris-HCl (pH 8), 100 mM NaCl, 10 μ M quinol, 10 μ M quinone, and 20 μ M oxidized cytochrome c. Only a statistically significant amount of superoxide production is shown as a percentage *above* the *bars*. *Error bars* represent S.D. of the mean of 12 measurements. DBH_2 , reduced form of 2,3-dimethoxy-5-methyl-6-decyl-1,4-benzoquinone.

mutant alleviates the effects of G167P to the point that $in\ vivo$ functionality of cytochrome bc_1 is restored. However, as the original 1Ala and 2Ala mutants affect the motion of ISP-HD to various degrees (10), the overall effects of a combination of G167P with 1Ala or 2Ala (G167P/1Ala or G167P/2Ala, respectively) differed. G167P/1Ala displayed generally higher activities and better growth than G167P/2Ala, which in terms of measured $in\ vitro$ activities was only slightly better than G167P and consequently displayed rather weak photosynthetic growth (compared with G167P/1Ala or WT) (Table 1 and Fig. 3).

EPR analysis revealed that the average position of ISP-HD in G167P/2Ala and G167P/1Ala sets at a longer distance from heme $b_{\rm L}$ than in the respective 2Ala and 1Ala mutants but still shorter than distances in G167P and WT (Fig. 5B). A new average position of ISP-HD for each double mutant (falling between the two average positions of ISP-HD in single mutants) indicates that the effects of two oppositely acting mutations (G167P versus 1Ala or 2Ala) add. The new positions extend the array of ISP-HD positions available for functional/structural studies.

Based on the results obtained with G167P and with double mutants G167P/2Ala and G167P/1Ala, we anticipate that there are other mutations in cytochrome b likely to exert an effect similar to that of G167P. The prominent candidates are mutations in the *ef* loop region of cytochrome *b* (such as L286F) identified as suppressors of 1Ala mutation (52). The suppression effect would be analogical to the effect of combination of G167P with 1Ala described here. Although the ef loop is in a different region of cytochrome b than helix cd1/cd2, it also forms a part of the interaction site of cytochrome b with ISP-HD. Furthermore, the ef loop was proposed to form a barrier that ISP-HD needs to cross to move out of the Q_o site upon its large scale movement toward cytochrome c_1 and to move back to the site (52). The suppression mutation L286F was proposed to diminish the barrier, facilitating the movement of ISP-HD in the presence of 1Ala (52). Conceivably, this would shift the average position of ISP-HD toward positions more remote from the Q_0 site as in G167P. However, the influence of L286F on motion of ISP-HD is not as large as in G167P as indicated by

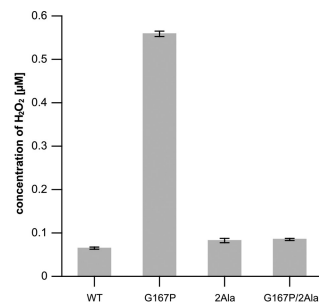


FIGURE 9. Concentration of $\rm H_2O_2$ accumulated during the course of the reaction catalyzed by WT cytochrome bc_1 and mutants (G167P, 2Ala, and G167P/2Ala) in the presence of SOD measured using the Amplex Red horseradish peroxidase method. Conditions were 50 mm Tris-HCl (pH 8), 100 mm NaCl, 10 μ M quinol, 10 μ M quinone, and 10 μ M oxidized cytochrome c from bovine heart. Error bars represent S.D. of the mean of four measurements.

less severe kinetic impediments of L286F in comparison with G167P and by the Ps^+ phenotype of the L286F mutant (as opposed to Ps^- G167P) (52). Another indication that the effects of G167P might be more profound than those of L286F comes from the observation that only G167P was able to suppress the effects of 2Ala.

G167P offered an important set of new conditions for investigating the mechanism of superoxide production by cytochrome bc_1 . According to a recently proposed mechanism, the back electron transfer from reduced heme b_1 to quinone (24, 25) results in formation of semiquinone at the Q_o site that has the best chance to react with oxygen when the ISP-HD occupies positions remote from the Q_o site at the time of semiquinone formation. The kinetic constraint associated with the position of ISP-HD was originally deduced from the observation that mutants arresting ISP-HD at the Qo site (2Ala and 1Ala) suppress ROS production by cytochrome bc_1 (25, 26). This model predicts that mutations that act in the opposite direction, i.e. shift the equilibrium position of ISP-HD toward a position more remote from the Q₀ site, should increase the level of ROS production. The results obtained with G167P are consistent with this prediction. Antimycin-inhibited G167P showed an increased level of superoxide production in comparison with antimycin-inhibited native enzyme. Furthermore, G167P generated superoxide even without any inhibitor present when the reaction mixture contained quinone in addition to quinol in the reaction mixture (Figs. 8 and 9). This was in contrast to the uninhibited native enzyme, 2Ala mutant, and G167P/2Ala double mutant, which did not generate detectable superoxide under any of the tested conditions (i.e. even with quinone present) in the measurements based on superoxide dismutase-sensitive reduction of cytochrome *c*.

The observation that it was necessary to add quinone to observe the generation of superoxide in uninhibited G167P is particularly interesting as it provides additional support for the notion that the initial reaction in the sequence leading to superoxide production is the electron transfer from heme $b_{\rm L}$ to quinone to form semiquinone in the ${\rm Q_o}$ site (24, 25). In this reaction, quinone acts as a substrate, explaining the negative slope of Fig. 8C (decrease in the level of superoxide generation with an increase in the quinol/quinone pool ratio). We note that it would be difficult to explain this slope on the basis of an alternative model assuming that the superoxide-generating semiquinone is formed upon one-electron reduction of Rieske cluster by quinol (in this case quinol, not quinone, is the substrate) (20, 21, 53, 54).

The conditions identified here under which uninhibited G167P generated superoxide are consistent with the observation reported earlier that the maximum rate of superoxide generation by complex III in mitochondrial system occurred when the Q pool was partly oxidized (24, 55). In fact, this finding was taken as initial evidence for the heme b_1 to quinone reaction at the Q_o site (24). In those studies as in the majority of other studies, the electron flow through cytochrome bc_1 must have been inhibited by antimycin to observe superoxide production. In this context, G167P shows that certain mutations in cytochrome b (especially those that promote ISP-HD to occupy positions remote from the Qo site) may induce changes that enhance superoxide generation by cytochrome bc_1 to the point that it becomes detectable in the non-inhibited enzyme. However, in our in vitro experiments, this appeared to strongly depend on the quinone/quinol ratio. Relating this to physiological conditions implicates that the level of generation of superoxide by certain mutants may depend on the redox state of the Q pool with a general tendency that the level will increase as the pool becomes more oxidized. It should be noted that the mutants themselves may influence the redox state of the Q pool; however, predicting a direction of this change (whether the Q pool becomes more oxidized or more reduced) is difficult. This is because such a change will be an overall effect of not just decreased enzymatic activity of the mutants (a typical effect also seen in G167P) but other factors as well (respiration rate, activity of other respiratory complexes, and membrane potential).

Remarkably, \sim 70% of ubiquinol identified as the condition associated with the maximum level of superoxide production in G167P (Fig. 8*B*) closely resembles the condition of maximal superoxide production reported for antimycin-inhibited submitochondrial particles (24). This shows that modulation of superoxide production by changes in the quinol/quinone ratio can be similar in both mitochondrial and bacterial complexes irrespective of the initial cause that led to the impediment in electron transfer (mutation or inhibitor). In higher organisms, this modulation could be a part of redox signaling with the Q_o site acting as a sensor of the redox state of the Q pool in membrane (1, 55, 56).

S151P (G167P in *R. capsulatus*) was originally reported in a patient with exercise intolerance (35). It was restricted to muscle tissue (80% heteroplasmy) with extremely decreased complex III enzymatic activity (35). Analysis of mitochondrial par-



ticles obtained from these cells indicated a slight decrease in the amounts of cytochrome b and cytochrome c_1 subunits (35). Conversely, the analogous mutation studied in a yeast model system resulted in a dramatic decrease in the level of ISP subunit and consequently in the enzymatic activity of the mutated complexes (34). Our results in a bacterial model system indicate that Pro at position 167 (151 in the human mitochondrial cytochrome b) causes a shift in the average position of ISP-HD toward positions more remote from the Q_o site. We propose that this effect could decrease the stability of complex III during biochemical preparations, for example by making it more susceptible to proteolytic digestion, which would in part account for the reduced amounts of various subunits observed in human tissues and the yeast model.

Interestingly, it was reported that in yeast the effect of S152P can be suppressed by a mutation, A90D (34), located in the hinge region of the ISP subunit. A90D in yeast is in the same region as Ala insertions in R. capsulatus (10) and thus may influence the motion of ISP-HD in analogy to the effects of 1Ala or 2Ala mutant. The observation that the effects of S152P (G167P) can be suppressed by the mutations located in the hinge region of the ISP subunit in both yeast and bacterial cytochrome bc_1 comes as additional support for the notion that the original mutation influences the movement of ISP-HD, resulting in the change of its average position.

We anticipate that there might be other human disease-related mutations in cytochrome b at the cytochrome b/ISP-HD interface that cause a shift in the average position of ISP-HD in a similar direction as G167P (ISP-HD more remote from the Q_o site). Such mutations are likely to influence the superoxidegenerating activity of complex III in a manner similar to G167P. Quite importantly, as G167P indicates here, this activity strongly depends on the redox state of the quinone pool (quinone/quinol ratio) and therefore may dynamically change in living cells.

Author Contributions—A. B. designed research and performed optical measurements (spectra, enzymatic activities, and ROS production), photosynthetic growth experiments, analyzed and interpreted the data, and assisted in writing the paper. P. K. performed CW EPR measurements and light-induced electron transfer, analyzed and interpreted the data, and assisted in writing the paper; R. E. performed molecular biology and gel analysis and contributed to photosynthetic growth experiments. R. P. contributed to CW EPR measurements and performed statistical analysis. M. S. performed pulse EPR measurements and contributed to analysis and interpretation of the data. A. O. designed research, analyzed and interpreted the data, and wrote the paper. All authors reviewed the results and approved the final version of the manuscript.

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