



## Role of Cooperative $H^+/e^-$ Linkage (Redox Bohr Effect) at Heme $a/Cu_A$ and Heme $a_3/Cu_B$ in the Proton Pump of Cytochrome $c$ Oxidase

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**Abstract**—It is a pleasure to contribute to the special issue published in honor of Vladimir Skulachev, a distinguished scientist who greatly contributes to maintain a high standard of biochemical research in Russia. A more particular reason can be found in his work (Artzabanov, V. Y., Konstantinov, A. A., and Skulachev, V. P. (1978) *FEBS Lett.*, **87**, 180-185), where observations anticipating some ideas presented in my article were reported. Cytochrome  $c$  oxidase exhibits protonmotive, redox linked allosteric cooperativity. Experimental observations on soluble bovine cytochrome  $c$  oxidase are presented showing that oxido-reduction of heme  $a/Cu_A$  and heme  $a_3/Cu_B$  is linked to deprotonation/protonation of two clusters of protolytic groups,  $A_1$  and  $A_2$ , respectively. This cooperative linkage (redox Bohr effect) results in the translocation of 1  $H^+$ /oxidase molecule upon oxido-reduction of heme  $a/Cu_A$  and heme  $a_3/Cu_B$ , respectively. Results on liposome-reconstituted oxidase show that upon oxidation of heme  $a/Cu_A$  and heme  $a_3/Cu_B$  protons from  $A_1$  and  $A_2$  are released in the outer aqueous phase.  $A_1$  but not  $A_2$  appears to take up protons from the inner aqueous space upon reduction of the respective redox center. A cooperative model is presented in which the  $A_1$  and  $A_2$  clusters, operating in close sequence, constitute together the gate of the proton pump in cytochrome  $c$  oxidase.

**Key words:** cytochrome  $c$  oxidase, proton pump, cooperative coupling

Allosteric cooperativity is a fundamental property in different hemoproteins and enzymes [1, 2]. This property confers to the protein the capacity to feel and respond to the physiological state of cells and overall organisms. The prototype of an allosteric protein is provided by hemoglobin. In hemoglobin the binding of  $O_2$  to the heme iron pulls this by 0.5 Å in the plane of the porphyrin ring [3]. This small movement of the iron, which pulls after it the histidine coordinated at the fifth position, is amplified by a conformational wave that modifies the surface contacts of the globin chains. The result is a sequential increase in the binding affinity from the first to the fourth oxygen molecule [3]. Another important conformational change initiated by oxygen binding is the rupture at the surface of the  $\beta$  globins of a salt bridge between aspartate 94 and histidine 146. As a consequence the  $pK$  of the histidine decreases with proton release in the aqueous phase. This cooperative interaction, known as the alkaline Bohr effect, since it takes place at pH above 6, makes the oxygen affinity of hemoglobin increase with pH [3, 4]. The

Bohr effect results in an amplified physiological function in that it makes hemoglobin bind oxygen in the lung where the pH can be slightly higher than 7.4 and to release oxygen to tissues where pH is around pH 7.4 or lower [2].

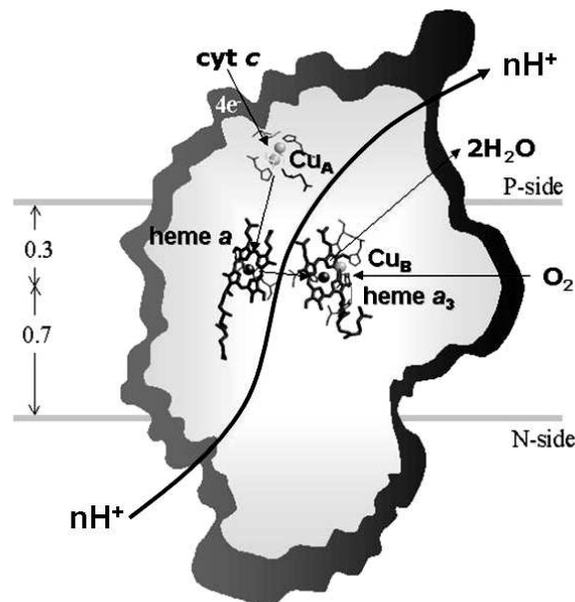
As far as membrane bound hemoproteins are concerned,  $b$  and  $a$  type cytochromes of energy-transfer respiratory chains exhibit a pH dependence of the midpoint redox potential [5]. It means that electron transfer at the metal is linked to proton exchange by protolytic groups in the enzyme [6]. Starting from this property of cytochromes, I proposed in the seventies that cooperative linkage between the redox state of the metal and  $pK$  of protolytic group(s) in the protein could result in redox-driven proton pumping across the coupling membrane if electron delivery to the metal was associated with  $pK$  increase of a group with proton uptake from one of the two aqueous phases separated by the osmotic barrier of the membrane, and electron transfer to a downhill redox center to  $pK$  decrease of the same group, or another in

proton connection with the first, with proton release in the opposite aqueous phase [7, 8]. This thermodynamic coupling between electron transfer and proton pumping, against a respiratory steady-state protonmotive force of some 250 mV in coupling membranes, will require an effective  $\Delta pK$  of 4–5 units and a  $\Delta E_h$  of 250–300 mV at least. It can be noted that this principle of vectorially organized redox-linked  $pK$  shifts of protolytic groups is now incorporated in some of the more recent models proposed for the proton pumping activity of cytochrome *c* oxidase [9–12]. It is feasible that cooperative protonmotive mechanisms may well operate in proton pumps in series or in parallel with direct primary protonmotive catalytic steps. An analogy can also be drawn, as anticipated some time ago [13], with the detailed models now put forward for the light driven proton pump of bacteriorhodopsin [14].

### REDOX-LINKED CONFORMATIONAL CHANGES IN CYTOCHROME *c* OXIDASE

Cytochrome *c* oxidase (Fig. 1) presents definite signs of redox-linked allosteric cooperativity. The  $E_m$  of the four metal centers of the oxidase ( $Cu_A$ , heme *a*, heme  $a_3$ , and  $Cu_B$ ) is pH dependent [5, 15–17]. There are anticooperative redox interactions of the metal centers [15, 18]. X-Ray crystallographic analysis shows redox-coupled structural changes in subunit I of the oxidase [19]. Picosecond time resolved tryptophan fluorescence also indicates redox-linked conformational changes [20]. In both cases the conformational changes appear to take place in the environment of heme *a*.

Non-resonant Raman spectroscopy has revealed in the oxidized bovine cytochrome *c* oxidase a peak at around  $1645\text{ cm}^{-1}$  that disappears in the reduced state [21]. Comparison of the Raman spectra of the unliganded oxidase with those exhibited by the CN-ligated oxidase, quinol oxidase which lacks  $Cu_A$ , cytochrome *c*, and imidazole derivatives of the extracted hemochrome allowed to assign the redox-sensitive Raman peak to a transition in the stretching vibrational mode of the C=C or C=N bond in the imidazole ring of one of the two axial histidine ligands of the low spin heme *a* [21]. Inspection of the X-ray crystallographic structures of the oxidized and fully reduced bovine oxidase reveals that reduction results in a few degree rotation of the imidazole plane of I-His378 on the axis perpendicular to the porphyrin plane [19] (see color insert, Fig. 2a). This movement of the histidine ligand, in a sense reminiscent of the movement of the histidine ligand induced by oxygen binding in hemoglobin, although small appears to be the closest sign of a perturbation of the protein structure induced by a change in the redox state of the heme *a* iron. The X-ray crystallographic structures show that the reduction of the oxidase results also in more significant conformational changes at



**Fig. 1.** Sketch of cytochrome *c* oxidase with location of the redox centers relative to the plane of the coupling membrane as obtained from X-ray crystallographic structures (see [10] for review).

heme *a*. Reduction causes the rupture of a hydrogen bond between the OH group of the hydroxyfarnesyl substituent of heme *a* and I-Ser382, with concomitant movement of serine and the hydrocarbon chain of the hydroxyfarnesyl [19]. A conformational wave induced by reduction of the oxidase appears to reach the cytosolic surface of subunit I in contact with subunit II. A segment of subunit I from I-Gly49 to I-Asn55 moves towards the surface with the carboxylic group of Asp51 becoming exposed to the aqueous phase [12, 19] (see color insert, Fig. 2b). Again an analogy seems to exist between these redox linked local/global conformational changes in the oxidase and those underlying the Bohr effect in hemoglobin.

The X-ray crystallographic structures show also a hydrogen bond between I-Arg38 and the formyl substituent of heme *a*, which was anticipated by resonance Raman spectroscopy studies [22] (Fig. 2b). This approach also revealed the presence of a water molecule near the formyl [9].

### REDOX-LINKED PROTON TRANSLOCATION IN CYTOCHROME *c* OXIDASE

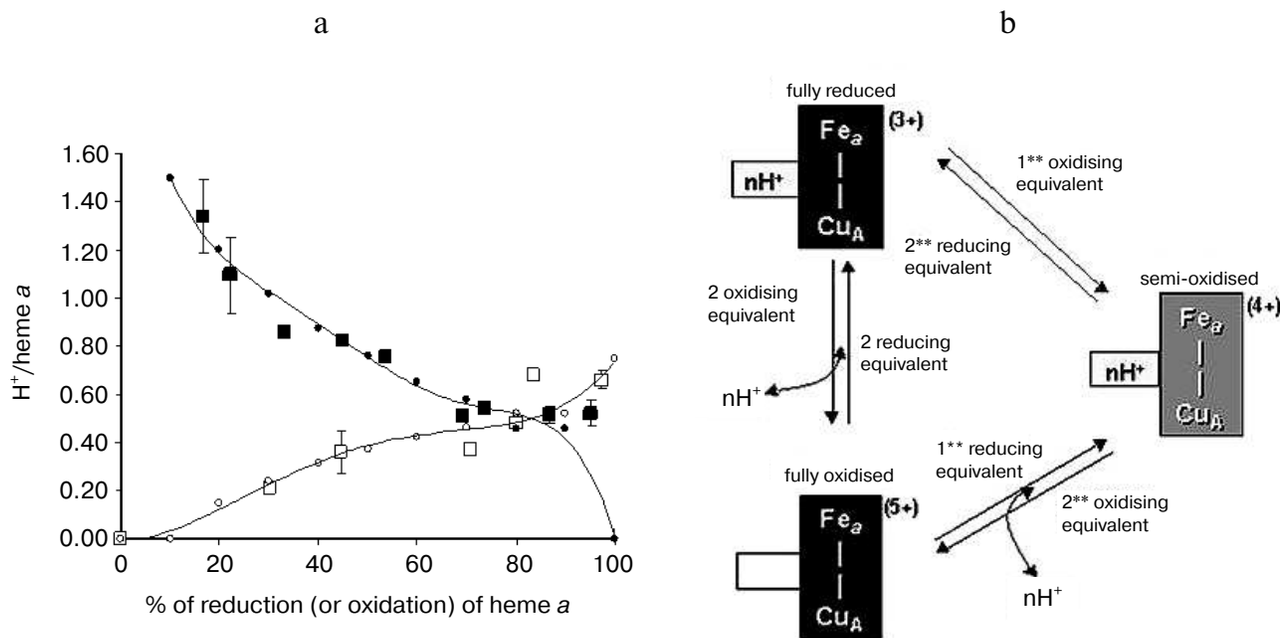
Measurements of proton release and uptake, associated with the oxidation and reduction respectively of heme *a* and  $Cu_A$  in the soluble CO-inhibited bovine cytochrome *c* oxidase (COX) ( $a_3$  and  $Cu_B$  clamped in the reduced state) have shown that oxido-reduction of these centers is coupled to  $H^+$  transfer, with an  $H^+/COX$  ratio

varying between 0.7 and 0.9 in the pH range 6.0-8.5 [23]. These data have been recently confirmed by Forte et al. who showed that in the soluble bovine [24] and *P. denitrificans* cytochrome *c* oxidase [25] the reduction of heme *a* is associated with the uptake of 0.6-0.8 H<sup>+</sup>/COX.

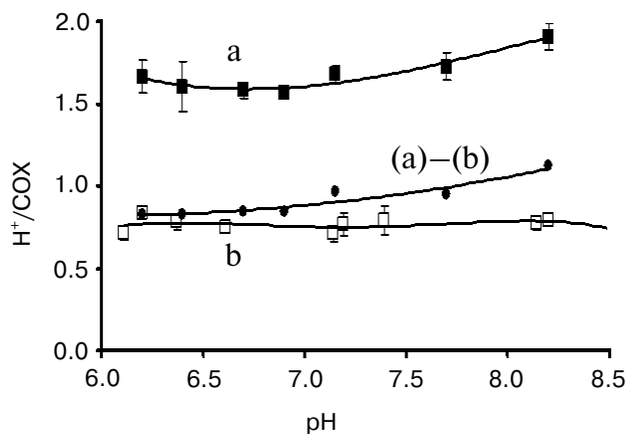
Electron/proton coupling at heme *a* only, with a H<sup>+</sup>/COX coupling ratio of 0.7-0.9, was apparently inconsistent with previous measurements of the pH dependence of the E<sub>m</sub> of heme *a*, which in the CO-inhibited COX was reported to amount to not more than -20 mV per pH unit increase in the same pH range [17]. A solution to this was provided by the finding that Cu<sub>A</sub> exhibits E<sub>m</sub> value and a pH dependence completely superimposed on that of heme *a*, the E<sub>m</sub> of both centers decreasing by around 20 mV/pH unit increase [23]. This shows that oxido-reduction of both heme *a* and Cu<sub>A</sub> is linked to p*K* shifts of two or more common acid-base groups, whose overall balance results in the observed H<sup>+</sup> release upon oxidation of the two centers and proton uptake upon their reduction [6]. The observed movement, upon reduction, of I-Asp51 from a hydrophobic interior to an aqueous environment, would result in proton dissociation from the carboxylic group of this residue [12, 19]. In the oxido-reduction of heme *a* this negative Bohr effect of I-Asp51 can however be associated with positive Bohr effects [26,

27]. Mathematical analysis of the pH dependence of proton transfer coupled to oxido-reduction of heme *a* and Cu<sub>A</sub> and of the E<sub>m</sub> of the two redox centers in the CO-liganded soluble cytochrome *c* oxidase resulted in a best-fit of the experimental points by an equation representing the case in which both heme *a* and Cu<sub>A</sub> shared coupling with a negative Bohr effect (I-Asp51, ?), p*K*<sub>ox</sub> 7.3 - p*K*<sub>red</sub> 4.0, and a minimum of two positive Bohr effects (p*K*<sub>ox</sub> 4.0 - p*K*<sub>red</sub> 6.9 and p*K*<sub>ox</sub> 5.4 - p*K*<sub>red</sub> 9.0, respectively) [27]. It seems significant in this context that S205 of subunit II, which is not too distant from the binuclear Cu<sub>A</sub> center, is in the crystal of the oxidized oxidase hydrogen bonded to I-Asp51 [12, 19]. In any event, it should be kept in mind that, in principle, the experimental points could be fitted by similar equations incorporating protolytic groups in the cluster with different p*K* shifts (cf. [28]).

An important consequence of the interactive coupling of the oxido-reduction of both heme *a* and Cu<sub>A</sub> with p*K* shifts in a common cluster of protolytic groups (A<sub>1</sub>) is that, while one electron reduction of the heme *a*/Cu<sub>A</sub> center is sufficient to produce maximal protonation of the cluster, release of the proton bound to the cluster will take place only when both heme *a* and Cu<sub>A</sub> are oxidized [29] (Fig. 3). The consequence is that at the steady state one electron at once has to pass through Cu<sub>A</sub> and heme *a* to



**Fig. 3.** Proton release/uptake coupled with oxidation/reduction of heme *a* and Cu<sub>A</sub> in CO-inhibited cytochrome *c* oxidase [29]. a) H<sup>+</sup> uptake/release associated with stepwise reductive-oxidative titration of Cu<sub>A</sub> and heme *a* in CO-liganded cytochrome *c* oxidase. The measured H<sup>+</sup>/heme *a*, Cu<sub>A</sub> ratios for proton uptake-release associated with consecutive stepwise reductive-oxidative titrations with hexammineruthenium (II) and ferricyanide are shown by filled and empty squares, respectively. Simulation of the H<sup>+</sup>/heme *a*, Cu<sub>A</sub> ratios is shown by small black circles for reductive titration and small empty circles for oxidation titration. The bars on the symbols (where given) indicate the standard errors of the mean value of three to four measurements. For experimental details see [29]. b) Model of interactive proton release/uptake coupled with oxidation/reduction of Cu<sub>A</sub> and heme *a* in CO-inhibited cytochrome *c* oxidase. Reproduced with permission from [29].



**Fig. 4.** pH dependence of proton release/uptake associated with oxidation/reduction of metal centers induced by ferricyanide/ferrocyanide in anaerobic soluble bovine heart cytochrome *c* oxidase. Filled squares (a):  $H^+/COX$  ratios associated with oxido-reduction of hemes *a*,  $a_3$ ,  $Cu_A$ , and  $Cu_B$  in the unliganded oxidase. Empty squares (b):  $H^+/COX$  ratios associated with oxido-reduction of heme *a* and  $Cu_A$  in the CO-liganded COX, heme  $a_3$  and  $Cu_B$  clamped in the reduced state [23]. Soluble cytochrome *c* oxidase was preincubated in 150 mM KCl before induction of the redox transitions. For calculation of the  $H^+/COX$  ratios in the unliganded oxidase presented in (a), molar extinction coefficient of  $0.188 \mu M^{-1} \cdot cm^{-1}$  [35, 36] for heme *a* plus heme  $a_3$  was used. This is higher than molar extinction coefficient of  $0.152 \mu M^{-1} \cdot cm^{-1}$  used in previous calculations [31, 33].

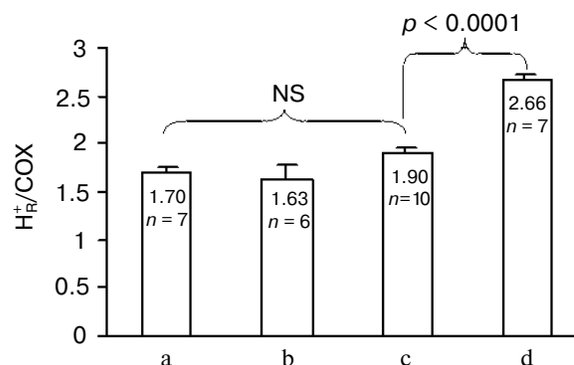
result in the translocation of around 1  $H^+$  per electron. This rule of “one electron at once” might represent one of the causes of the slip, observed under certain conditions, in the proton pump of cytochrome *c* oxidase [27, 30, 31].

The vectorial nature of Bohr protons linked to oxido-reductions of heme  $a/Cu_A$  was investigated in the mixed valence CO-liganded bovine oxidase reconstituted in liposomes, in which heme  $a_3$  and  $Cu_B$  at the binuclear site were clamped in the reduced state [29]. The oxidation of heme  $a/Cu_A$ , reduced by an external reductant, was associated with  $H^+$  release in the external aqueous phase. The expected  $H^+$  uptake associated with re-reduction of heme  $a/Cu_A$  was considerably delayed with respect to completion of the reduction of these centers, unless CCCP was added [29]. These results thus showed that whilst oxidation of heme  $a/Cu_A$  results in the release in the external bulk phase of the Bohr protons linked to these centers, their reduction results in the uptake of the Bohr protons from the inner space. Consistent with this picture are previous results of Artzabanov et al. [32] who showed oxidation/reduction of heme *a* upon alkalization/acidification, respectively, of the matrix space in intact rat liver mitochondria.

Redox Bohr protons have also been analyzed in the soluble unliganded bovine cytochrome *c* oxidase. In this case the fully oxidized oxidase was preincubated in the presence of 150 mM KCl, conditions under which the

binuclear center is reported to be occupied by  $Cl^-$ , thus preventing  $OH^-/H_2O$  exchange at this site [24]. Oxido-reduction of the four metal centers in the unliganded oxidase resulted in the release/uptake of 1.7-1.9  $H^+/COX$ . This  $H^+$  exchange is, at pH values around neutrality, around twice that observed upon oxido-reduction of heme  $a/Cu_A$  in the mixed-valence CO-liganded oxidase (Fig. 4) [31, 33]. Thus, oxido-reduction of heme  $a_3/Cu_B$  at the binuclear center results in the exchange at a second cluster of protolytic groups,  $A_2$ , of 0.8-0.9 Bohr protons.

The contribution of cooperative  $H^+/e^-$  linkage at heme  $a/Cu_A$  and  $a_3/Cu_B$  (redox Bohr effects) in proton pumping was investigated in bovine cytochrome *c* oxidase reconstituted in liposomes. In a series of experiments the four metal centers in the unliganded oxidase were fully reduced in anaerobiosis by the photo activated EDTA-riboflavin system [34]. The four centers were then oxidized by a stoichiometric amount of ferricyanide. This resulted in a synchronous  $H^+$  release in the external bulk phase that amounted to an  $H^+/COX$  ratio of  $1.63 \pm 0.37$  (Fig. 5).



**Fig. 5.** Proton release associated with anaerobic ferricyanide oxidation of unliganded fully reduced soluble (a) and liposome-reconstituted bovine cytochrome *c* oxidase (b), aerobic oxidation (c) and aerobic oxidation-re-reduction (d) of liposome-reconstituted cytochrome *c* oxidase. a) Anaerobic reduction of cytochrome *c* oxidase ( $1 \mu M aa_3$  plus  $0.5 \mu M$  cytochrome *c*) was attained by succinate oxidation in the presence of a trace of frozen-thawed broken beef-heart mitochondria. Oxidation was produced by the addition of an amount of ferricyanide stoichiometric with the sum of the reduced metal centers in the presence of antimycin plus myxothiazol [33]. b, c) Anaerobic full reduction of the oxidized oxidase,  $0.8-1.0 \mu M aa_3$  plus  $0.5 \mu M$  cytochrome *c* (b),  $0.5-1.0 \mu M aa_3$  plus  $0.1 \mu M$  cytochrome *c* (c) was attained by the photo-activated EDTA/riboflavin system [34]. d) Anaerobic full reduction of the oxidase,  $1 \mu M aa_3$  plus  $1 \mu M$  cytochrome *c*, was produced by 1 mM ascorbate and  $20 \mu M$  hexammineruthenium. Oxidation was produced by the addition of an amount of ferricyanide stoichiometric with the sum of the reduced metal centers (b), or with an amount of  $O_2$  stoichiometric with that of  $aa_3$  (c and d). The concentration of  $aa_3$  was determined using molar extinction coefficient of  $0.188 \mu M^{-1} \cdot cm^{-1}$  for heme *a* plus heme  $a_3$  [35, 36]. This is higher than molar extinction coefficient of  $0.152 \mu M^{-1} \cdot cm^{-1}$  used in previous calculations [27]. (Capitanio, G., Martino, L., Capitanio, N., and Papa, S., unpublished results.)

In another series of experiments the reconstituted, unliganded oxidase, fully reduced in anaerobiosis, was subjected to oxidation by a stoichiometric amount of oxygen, in the absence or presence of ferrocyclochrome *c*. The aerobic oxidation of the oxidase resulted in fast  $H^+$  release in the external phase amounting, in a number of these experiments, to an  $H^+/COX$  ratio of  $1.9 \pm 0.06$  (Fig. 5). When the oxidation was immediately followed by a complete re-reduction of the metal centers of the oxidase by ferrocyclochrome *c*, an additional  $H^+$  release was observed, which brought the overall  $H^+/COX$  ratio to  $2.66 \pm 0.07$  (Fig. 5). It should be noted that for similar experiments Verkhovsky et al. [36] reported an  $H^+/COX$  ratio for the oxidation phase approaching 2, thus practically equal to that measured by us, but an overall  $H^+/COX$  ratio approaching 4 when the oxidation phase was immediately followed by re-reduction of the metal centers. The reasons for this latter discrepancy are at present not clear (see also [37]).

The  $H^+/COX$  ratio for proton release in the external aqueous phase associated with the oxidation of the metal centers by ferricyanide in the reconstituted oxidase was practically equal to the  $H^+/COX$  ratio for proton release induced by ferricyanide oxidation of the metal centers in the soluble oxidase. The  $H^+/COX$  ratio for proton release induced by aerobic oxidation of the reconstituted oxidase was also very close, if not equal, to the above  $H^+/COX$  ratios. It has to be concluded that the  $H^+$  release observed in the ferricyanide and the  $O_2$  oxidation of the fully reduced reconstituted oxidase derives from deprotonation of protolytic groups in the clusters  $A_1$  and  $A_2$  responsible for the redox Bohr effects linked to heme *a*/ $Cu_A$  and heme  $a_3$ / $Cu_B$ , respectively.

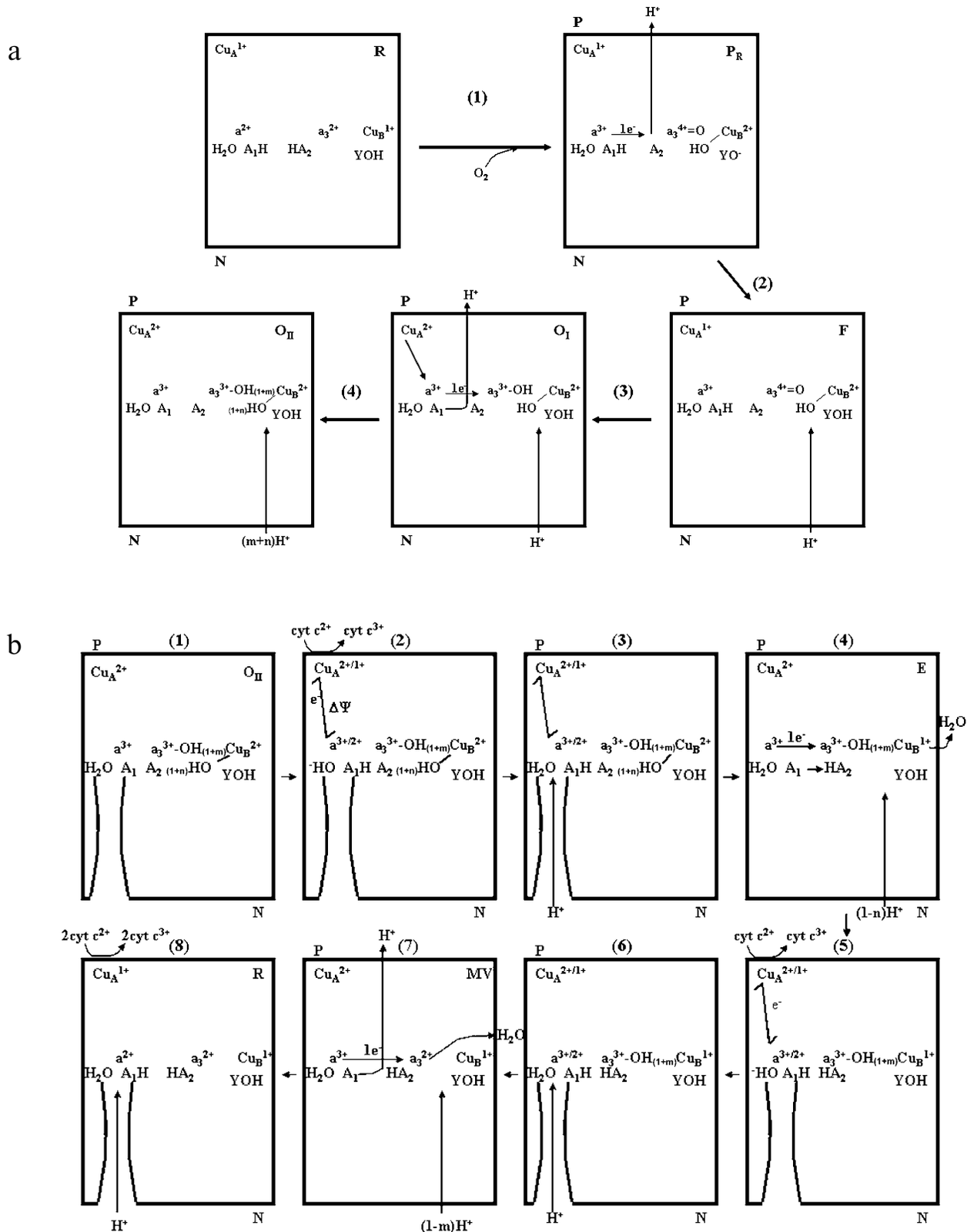
$H^+/COX$  ratios obtained from simulations based on different membrane orientation of the Bohr effect linked to oxido-reduction of heme  $a_3$ / $Cu_B$  and  $H^+$  pumping in the normal or reverse direction associated to reverse electron flow in the ferricyanide oxidation of reduced oxidase were found to be incompatible with the experimental  $H^+/COX$  ratios.

## DISCUSSION

Schemes on Figs. 6a and 6b describe the sequence of electron and proton transfer steps in a single turnover, in which the fully reduced enzyme is oxidized by oxygen and then re-reduced by ferrocyclochrome  $c^{2+}$ . In the fully reduced oxidase both the clusters  $A_1$  and  $A_2$  associated to heme *a*/ $Cu_A$  and heme  $a_3$ / $Cu_B$  respectively are in the protonated state (Fig. 6a).  $A_1$  and  $A_2$  each bind approximately one  $H^+$ . The number of these protons appears, however, to vary slightly with pH [23, 31, 33]. Upon binding of  $O_2$  (intermediate A is not shown) with oxidation of heme *a*, heme  $a_3$ , and  $Cu_B$  in the reductive cleavage of  $O_2$  to form the  $P_R$  compound the Bohr proton of the cluster  $A_2$

is released in the outer aqueous space (P).  $Cu_A$  (or heme *a*) being still reduced holds the Bohr proton on  $A_1$ . In the conversion of R to  $P_R$  no scalar proton is taken up from the inner aqueous space (N) [10, 38]. A tyrosine of subunit I, I-Tyr244 (bovine numbering) [19, 39, 40], or another tyrosine [41] connected with the binuclear center, donates the proton consumed in the formation of  $Cu_B^{2+}-OH$ . In the  $P_R \rightarrow F$  conversion, one proton is taken up from the inner space and reprotonates I-Tyr- $O^-$  [42]. F is converted to  $O_I$  upon reduction by  $Cu_A^{1+}$  of  $a_3^{4+}=O$  to  $a_3^{3+}-OH$ , with simultaneous uptake of one scalar  $H^+$  from the inner space [10, 37]. The oxidation of both  $Cu_A$  and heme *a* results in the release in the outer space of the Bohr proton of cluster  $A_1$ .  $O_I$  is then converted to  $O_{II}$  with partial protonation to  $H_2O$  of the two  $OH^-$  bound to  $a_3$  and  $Cu_B$ , the extent of this depending on the actual pH [31, 33]. In conclusion, in the oxidative phase of a single turnover two  $H^+$  deriving from the  $A_1$  and  $A_2$  clusters in the oxidase are released in the outer space. No proton pumping from the inner to the outer space would take place in this phase.

In the re-reduction of  $O_{II}$  (Fig. 6b) the first electron delivered to the oxidase rapidly distributes itself between  $Cu_A$  and heme *a*, this is associated with transfer of a proton from an  $H_2O$  near the formyl of heme *a* to the cluster  $A_1$  [9, 12]. In this phase the only electrogenic event is represented by the rapid 10–20  $\mu$ sec electron transfer from  $Cu_A$  to heme *a* located some distance below the outer surface (cf. [43–45]) (Fig. 1). The proton movement from  $H_2O$  to  $A_1$  might cover a short distance essentially parallel to the plane of the membrane, thus it will not result in significant charge translocation along the axis perpendicular to the plane of the membrane. The  $OH^-$  generated near heme *a* is then released in the outer space or neutralized by an incoming proton from the inner space. This phase, too, might not result in the generation of a transmembrane potential, if the  $H_2O/A_1$  site at heme *a* is exposed to a  $H_2O$  channel continuous with the outer aqueous bulk phase [12]. The single electron now moves from heme *a* to  $Cu_B$  (intermediate E) with electrogenic uptake from the inner phase of  $(1 - n)H^+$  consumed in the protonation of the  $OH^-$  to  $H_2O$  at  $Cu_B$ . The electron transfer from heme *a*/ $Cu_A$  to  $Cu_B$  results in proton transfer from the  $A_1$  cluster (whose effective p*K* decreases) to the  $A_2$  cluster (whose p*K* simultaneously increases). No proton release in the outer bulk phase would take place in the transfer of the first electron from heme *a*/ $Cu_A$  to heme  $a_3$ / $Cu_B$  (cf. [43–45]). Delivery of a second electron to  $Cu_A$ /heme *a* results again in proton transfer from  $H_2O$  to  $A_1$  at the heme *a*.  $OH^-$  is reprotonated to  $H_2O$  by  $H^+$  from the inner space. Transfer of the second  $e^-$  to heme  $a_3$  is associated with protonation of the second  $OH^-$  to  $H_2O$  at the binuclear site by  $(1 - m)H^+$  from the inner space. Oxidation of both heme *a* and  $Cu_A$  results in deprotonation of  $A_1$ . Since  $A_2$  is already protonated, an  $H^+$  is released in the bulk phase [46]. This situation might be



**Fig. 6.** Schemes describing the sequence of electron transfer and proton translocation steps in a single turnover transition of the membrane associated cytochrome *c* oxidase from the fully reduced (R) to the oxidized ( $O_{II}$ ) state (a) and from the  $O_{II}$  to the R state (b). For details, see text.

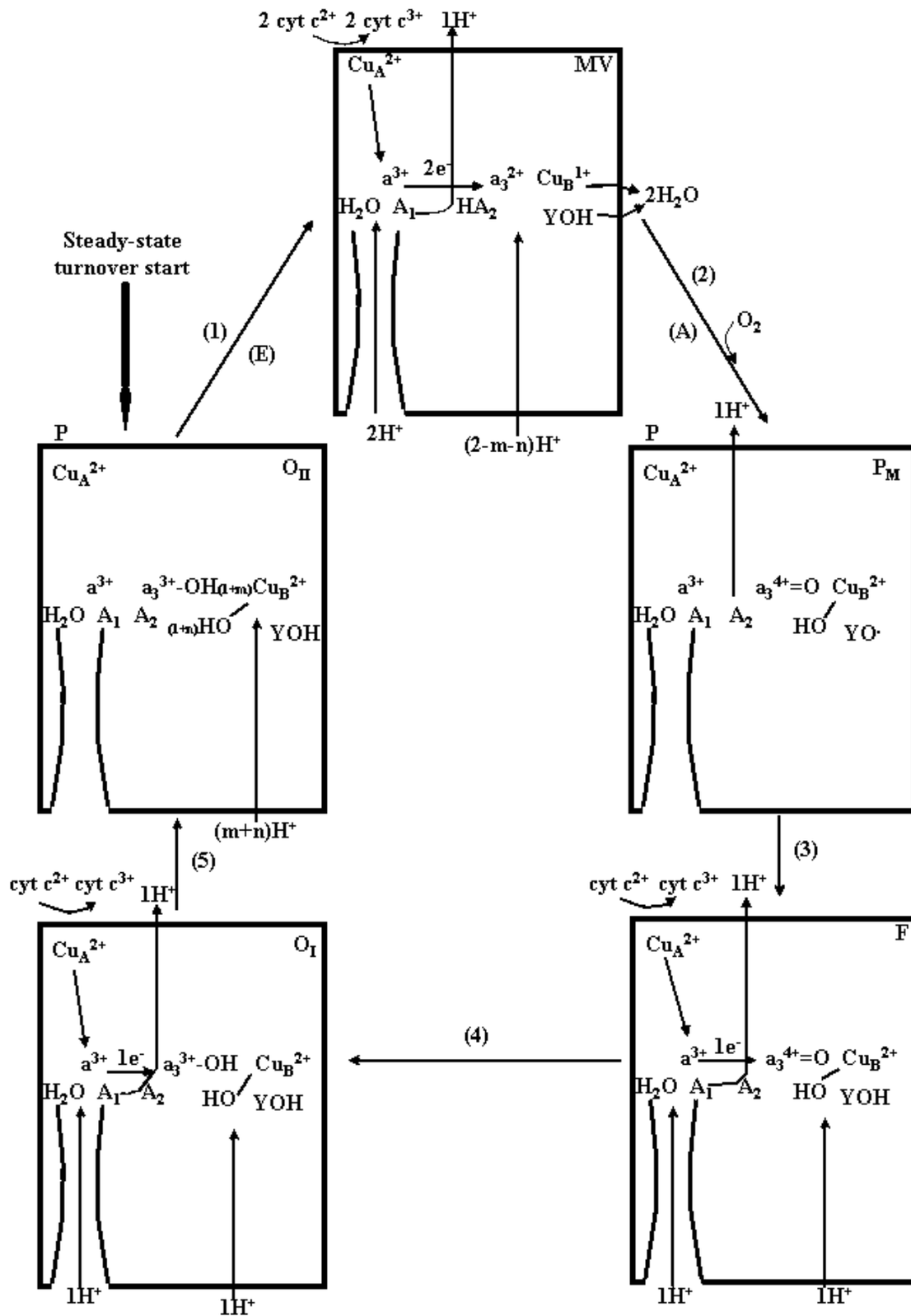


Fig 7. Catalytic cycle for reduction of  $O_2$  to  $H_2O$  by ferrocyanochrome  $c$  and proton pumping in the cytochrome  $c$  oxidase in the coupling membrane at the respiratory steady-state. For details, see text.

mimicked by the CO-inhibited oxidase in which heme  $a_3$  and  $\text{Cu}_B$  are clamped in the reduced state. Finally, delivery of two additional electrons converts the oxidase in the fully reduced state in which both  $A_1$  and  $A_2$  are protonated. Thus, one  $\text{H}^+$  is pumped from the inner to the outer space in the reductive phase of the single turnover of the enzyme. The schemes described in Fig. 6 (a and b) result in an  $\text{H}^+/\text{COX}$  ratio of 3 (see Fig. 5) for the overall  $\text{H}^+$  release in the outer bulk phase in the  $\text{R} \rightarrow \text{O}_{II} \rightarrow \text{R}$  transition. As mentioned above Verkhovsky et al. [36] have reported an overall  $\text{H}^+/\text{COX}$  ratio of 4 for this transition. A higher ratio would require additional  $\text{H}^+$  release at one of the steps described in the scheme of Fig. 6b and/or some aerobic re-oxidation of the metals at the binuclear site during the re-reduction phase.

The scheme in Fig. 7 describes the sequence of electron transfer and proton translocation steps in the catalytic cycle of cytochrome *c* oxidase at the respiratory steady state, resulting in pumping of 1  $\text{H}^+$  from the inner to the outer aqueous phase per each electron being transferred from ferrocycytochrome *c* to  $\text{O}_2$ . It should be noted that, at difference of the single turnover  $\text{R} \rightarrow \text{O}_{II} \rightarrow \text{R}$  transition, at the steady state one electron at once is transferred, through  $\text{Cu}_A/\text{heme } a$ , from ferrocycytochrome  $c^{2+}$  to the binuclear site. Furthermore, the fully reduced R compound is not formed during the aerobic steady-state catalytic cycle (see also [37]). This can explain difference in the  $\text{H}^+/\text{COX}$  ratios for overall proton release in the outer bulk phase in the  $\text{R} \rightarrow \text{O}_{II} \rightarrow \text{R}$  transition with respect to that in the reduction of  $\text{O}_2$  to  $\text{H}_2\text{O}$  in the steady-state catalytic cycle of the oxidase.

Under normal physiological conditions, a turnover of the oxidase starts with the fully oxidized enzyme. Reduction of the binuclear center by two electrons, delivered by ferrocycytochrome  $c^{2+}$ , one at once via  $\text{Cu}_A/\text{heme } a$  to  $a_3/\text{Cu}_B$ , results in the uptake of  $2\text{H}^+$  from the inner space. One  $\text{H}^+$  is held by the  $A_2$  cluster, whose effective  $\text{pK}_a$  increases upon reduction of  $a_3/\text{Cu}_B$ , the other  $\text{H}^+$  is pumped in the outer space.  $\text{O}_2$  binds to the reduced state (compound A is not shown) and upon reductive cleavage of  $\text{O}_2$  with oxidation of  $a_3/\text{Cu}_B$  (state  $\text{P}_M$ ) the second proton, taken up from the inner space in the previous reductive step and held on  $A_2$  [46], is pumped in the outer space. Note that, in the respiratory catalytic cycle for the reductive cleavage of  $\text{O}_2$ , three electrons derive from  $a_3/\text{Cu}_B$  reduced in the previous step, and a fourth electron with a proton is provided by I-Tyr [10, 37, 41]. In the transitions from  $\text{P}_M$  to F and F to  $\text{O}_I$  one electron at once is transferred from cytochrome *c*, via  $\text{Cu}_A/a$ , to the binuclear site with pumping of 1  $\text{H}^+$  from the inner to the outer space for each of the two steps. In the cycle the pumped protons follow a pathway whose critical elements are represented by the two clusters  $A_1$  and  $A_2$  both undergoing  $\text{pK}$  shift upon reduction/oxidation of the heme  $a/\text{Cu}_A$  and heme  $a_3/\text{Cu}_B$ , respectively. These clusters do not contribute scalar protons consumed in the reduction

of  $\text{O}_2$  to  $\text{H}_2\text{O}$  at the binuclear site. These protons are taken up from the inner space by a separate pathway.

There is a wealth of structural and functional observations showing that oxido-reduction of the low spin heme  $a/\text{Cu}_A$  is associated with protonmotive local/global cooperative conformational changes. The  $\text{H}^+/\text{e}^-$  coupling at heme  $a/\text{Cu}_A$  appears to represent the primary step in the uptake of pumped protons from the inner aqueous space. This proton uptake involves a cluster of protolytic groups  $A_1$ , whose effective  $\text{pK}$  is governed by the redox state of heme  $a/\text{Cu}_A$ . Protons taken up by cluster  $A_1$  upon reduction of  $\text{Cu}_A/\text{heme } a$  move, upon electron transfer from heme  $a$  to heme  $a_3$ , to cluster  $A_2$ , whose effective  $\text{pK}$  is governed by the redox state of  $a_3/\text{Cu}_B$ . Upon oxidation of  $a_3/\text{Cu}_B$  protons are released from  $A_2$  in the outer bulk phase. The cooperative  $\text{H}^+/\text{e}^-$  coupling at  $a_3/\text{Cu}_B$  synchronizes the pumping process with the partial steps of  $\text{O}_2$  reduction to  $\text{H}_2\text{O}$  and allows the energy, thus made available, to pump protons against a steady state protonmotive force of some 250 mV. The  $A_1$  and  $A_2$  clusters operating in close sequence constitute together the gate of the pump. The energy needed to pump protons from the inner to the outer space is distributed between  $\text{pK}$  shifts of  $A_1$  and  $A_2$  and is provided by the steps of  $\text{O}_2$  reduction to  $\text{H}_2\text{O}$  at the binuclear site. The  $A_1$ – $A_2$  pathway for proton pumping has to be separated from the pathway along which the scalar protons consumed in the reduction of  $\text{O}_2$  to  $\text{H}_2\text{O}$  are taken up from the inner space. This prevents pumped protons from being annihilated in the reduction of  $\text{O}_2$  to  $\text{H}_2\text{O}$ .

A discussion of the specific residues contributing to the  $A_1$  and  $A_2$  clusters, as well as the proton conduction pathways used for proton uptake from the inner space and release in the outer space, is beyond the scope of this paper. It can, however, be argued that I-Arg38 and the propionates of heme  $a$  contribute to cluster  $A_1$ , whilst the propionates of heme  $a_3$  and I-Arg438 and I-Arg439 contribute to cluster  $A_2$  [10, 37]. Development of the present working hypothesis is associated with clarification of these issues.

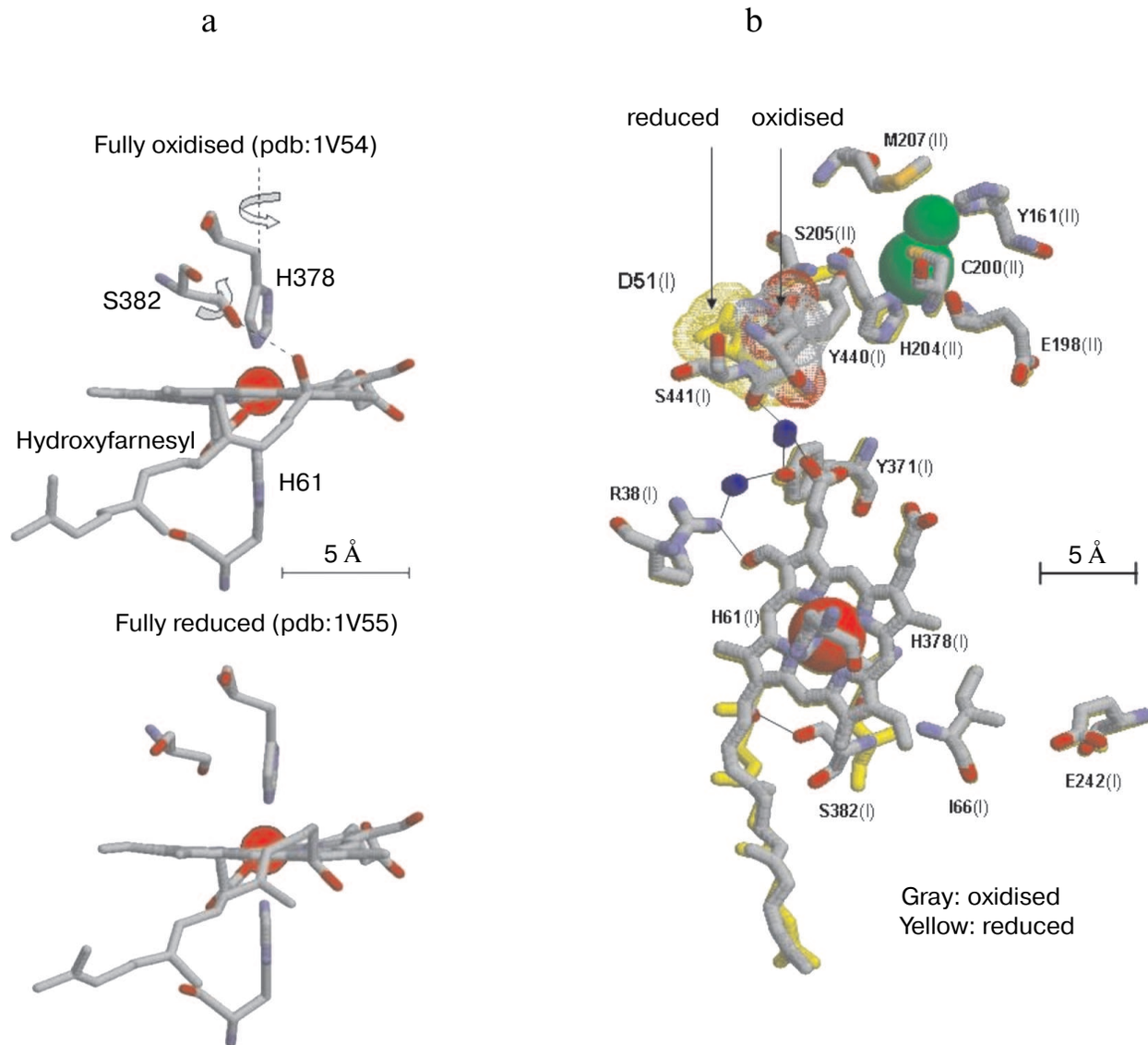
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**Fig. 2.** Redox-linked conformational changes in bovine cytochrome *c* oxidase. The picture was elaborated with the Rasmol 26 program using the 1V54 atomic coordinates from the PDB bank of the monomer of cytochrome *c* oxidase, superimposing the 1V54 (1.8 Å resolution) and the 1V55 (1.9 Å resolution) atomic coordinates from the PDB data bank of the oxidized and reduced crystal structure of bovine cytochrome *c* oxidase, respectively (“What if” program). a) Details of the redox-linked movement of I-H378, I-S382, and hydroxyl-farnesyl at the low spin heme *a* [19, 21]. b) The redox-linked movement of I-Asp51 (see text and [12, 19]) is evidenced by dotting in color van der Waals atomic radii. The stick structures of other residues shown evidenced only minor position changes of their atomic coordinates. The thin line shows possible hydrogen bond networks [12, 19].