Introduction

Cytochrome c oxidase (CcO; Ec 1.9.3.1) is the terminal enzyme of the inner mitochondrial electron transport chain (18). Bovine heart CcO is consisted of 13 different subunits (6) and the enzyme is almost always dimeric in three-dimensional lattice as it was obtained from crystallographic data (15).

The three largest subunits (I–III) are encoded by mitochondrial DNA and are synthesized in the mitochondria. They constitute the functional core of the enzyme and include all of the redox-active centers. The smaller subunits (IV–VIII) are nuclear encoded and most likely have a structural and/or regulatory role (4,5).

Treatment of CcO with physical (high hydrostatic pressure) or chemical (chaotropic agents, e.g., urea – 13, peroxidation by hydrogen peroxide – 9) factors enables to study the structural effects. Consequently, the relation between structural changes and enzymatic activity may be determined. We report the effects of hydrostatic pressure on electron transport activity and structural composition of CcO. Our approach is the study of pressure influence on CcO in view of different behavior of dimeric and monomeric CcO forms.

Materials and methods

Materials. Bovine cytochrome c oxidase was prepared from Keilin-Hartree heart particles by the method of Fowler et al. (2) with modifications described by Mahapatro and Robinson (8). The final oxidase pellet was solubilized in 100 mM phosphate buffer, pH 7.4 containing 1 % sodium cholate and 1 mM EDTA. Individual drops of purified enzyme (25 mg/mL protein) were quickly frozen by pipetting the solution into liquid nitrogen and stored at -80 °C. Cytochrome c oxidase concentrations were calculated on the basis of C–422 = 1.54 x 10^5 M^-1 cm^-1 (16). Horse heart cytochrome c (Type III) was purchased from Sigma Chemical Co. Reduced cytochrome c was freshly prepared by dithionite reduction, and an excess dithionite was removed by G-25 Sephadex gel filtration. Initial concentrations of ferrocytochrome c were determined using C–550 = 21 mM^-1 cm^-1. All other chemicals were reagent grade.

Methods. The high hydrostatic pressure experiments were performed as described in (11). Kinetics measurements and fluorescence spectra were recorded, respectively. In case of kinetics measurements tryptophan fluorescence was monitored (λ_ex = 295 nm, λ_em = 326 nm) as a function of time. Fluorescence spectra of tryptophan were recorded in emission range 300–500 nm using excitation wavelength 295 nm.

Determination of cytochrome c oxidase activity. Molecular activity of CcO was assayed spectrophotometrically at 25°C with 25–30 µM ferrocytochrome c as a substrate in 25 mM phosphate buffer, pH 7.0, containing 1 mg/mL decyl maltoside. Molecular activities were calculated by the pseudo-first-order rate of oxidation of ferrocytochrome c by high hydrostatic pressure.
1.75 nM cytochrome c oxidase (1). The activity of the purified enzyme was 300–350 s⁻¹ to an accuracy of ± 5%.

**HiTrapQ FPLC ion exchange chromatography.** Anion exchange chromatography on the HiTrapQ column was a modification of the method developed for a MonoQ FPLC column (7) and was used as described in (7).

**Analysis of subunits.** Nuclearily encoded subunits (subunits IV–VIII, nomenclature according to Kadenbach et al. (6) were determined by C₁₈ reversed phase HPLC (7). Absorbance intensity of subunit Vb was taken as the intensity standard for the chromatogram normalization. Determination of three mitochondrially encoded subunits (subunits I–III) was performed by SDS-polyacrylamide gel electrophoresis on 15% acrylamide gels that contained 2 M urea in addition to 0.1% SDS (12). Subunit content was determined by scanning on Personal Densitometer SI (Molecular Dynamics) using ImageQuant version 5.2 software.

**Results**

Monomeric and dimeric CcO were prepared by manipulating the concentrations of dodecyl maltoside and sodium cholate: high concentrations of dodecyl maltoside (5–10 mM) produce monomeric CcO while a mixture of dodecyl maltoside and sodium cholate produce dimeric CcO [10]. Homogeneity of the monomeric or dimeric forms was confirmed by sedimentation velocity as previously described (17).

**Effect of high pressure on the electron transport activity of cytochrome c oxidase.** Molecular activity of dimeric and monomeric CcO was assayed spectrophotometrically and obtained results showed a significant difference in the pressure-induced effect of dimer, and monomer, respectively. Exposure of dimeric CcO to pressure causes minimal loss of activity of enzyme (7–13% at the pressures 2.00–2.75 kbar) (Fig. 1). On the other hand the same pressures induce remarkable decrease in activity of monomeric CcO (24–48%) (Fig. 1). From these results we can conclude that dimeric CcO is functionally more stable than monomeric one if both forms are exposed to the equal hydrostatic pressure.

**Effect of high pressure on the structural composition of cytochrome c oxidase.** We used HiTrapQ ion exchange chromatography, combined with C₁₈ reversed-phase HPLC and SDS-PAGE for quantitatively assessing a subunit content of CcO. Chromatographic elution of purified CcO from a HiTrapQ FPLC ion exchange column allows a separation of intact 13-subunit CcO complex from a form that is missing subunits (7). Figure 2a shows HiTrapQ FPLC elution of dimeric CcO after exposure to high hydrostatic pressure 2.5 kbar. Protein eluted as the peak A and a small peak B without a significant difference between pressure-treated CcO and control sample (Fig. 2a). The gel electrophoresis and C₁₈ HPLC chromatography confirmed that peak A contains intact 13-subunit complex and peak B contains 11 subunits and is completely devoid of subunit VI and VIb.

**Fig. 1:** Dependence of dimeric and monomeric cytochrome c oxidase activity on high hydrostatic pressure. Dimeric CcO (5 µM) (circles) was prepared in 20 mM Tris-Cl pH 7.4 with 1mg/mg protein dodecyl maltoside in absence of dialysis. Monomeric CcO (5 µM) (squares) was prepared with 5mg/mg protein dodecyl maltoside followed by dialysis overnight in 20 mM Tris-Cl pH 7.4 with 0.1mg/mL dodecyl maltoside.

**Fig. 2:** HiTrapQ FPLC ion exchange chromatography elution of dimeric (a) and monomeric (b) cytochrome c oxidase as measured after exposure to high hydrostatic pressure. Both forms were prepared as described in Fig. 1. The elution gradient was as described previously (7). HiTrapQ elution of dimer (a) after exposure to 2.5 kbar is represented by trace 2. Trace 1 shows the elution of control sample. The elution of monomer (b) after exposure to 2.00 kbar is represented by trace 2, and to 2.5 kbar by trace 3. Trace 1 shows elution of sample in absence pressure.
Different situation became when the monomeric CcO was exposed to high hydrostatic pressure (Fig. 2b). Increase of pressure to 2.5 kbar caused a remarkable change in the chromatographic elution of monomeric CcO (Fig. 2b trace 3). As it can be seen from figure, the peak C was the most intensive and the peak B completely disappeared.

Reversed-phase C\textsubscript{18} HPLC enables to determine the 10 nuclear encoded subunits (IV – VIII), which are very difficult to separate by SDS – PAGE (7). We used this method for a determination of monomeric CcO subunit composition of eluting protein shown in Fig. 2b (peaks A, C) (Fig. 3).

The chromatographic analysis of the peak C shows a loss of subunits VI\textsubscript{a}, VI\textsubscript{b}, VII\textsubscript{a} (Fig. 3). Content of subunit VII\textsubscript{a} was only 17\% from that in the absence pressure and the intensity of subunits VI\textsubscript{a} (2\%) and VI\textsubscript{b} (8\%) decreased enormously (Fig. 3). Similar results were obtained when monomeric CcO was exposed to 2.75 kbar pressure. From our findings, we can conclude that the most of the monomeric CcO exposed to pressure 2.5 kbar and higher values occurs in the form, which is missing subunits VI\textsubscript{a}, VI\textsubscript{b} and VII\textsubscript{a} (Figures 2b and 3).

Determination and quantitation of the three largest subunits (I – III) was provided by SDS-PAGE. The composition of the monomeric CcO exposed to 2.5 kbar eluting in the HiTrap elution as peaks A, C (Fig. 2b, trace 3) as well as the control enzyme resulting in peaks A, B is shown in Fig. 4, columns 1–4.

The intact 13–subunit control CcO eluted as peak A is presented in column 1 in Fig. 4 and the intact 13–subunit complex of CcO exposed to 2.5 kbar is shown in column 2. Although we can see a small difference in intensity of subunit III band between control sample and sample exposed to pressure, the subunit III is 96\% the control sample. Remarkable change was recorded in case of peak C. Subunit III band of the pressure treated enzyme completely disappeared (Fig. 4, column 4) and the percent of this subunit is only 6\% the control CcO. From results obtained using SDS-PAGE it is clear that not only subunits VI\textsubscript{a}, VI\textsubscript{b}, VII\textsubscript{a} but also subunit III is missing in peak C elution of monomeric CcO. We can conclude: the exposure of monomeric CcO to pressure higher than 2.5 kbar leads to changes of its composition and therefore, disturbs a structural integrity of monomeric enzyme. With dimeric enzyme, only subunits VI\textsubscript{a} and VI\textsubscript{b} dissociate, neither of which is essential for full electron transport activity.

**Discussion**

High hydrostatic pressure was used as a physical factor for a study of possible different behavior of dimeric and monomeric CcO. Both forms of CcO were estimated from point of view of its partial functional stability (pressure effect on redox- state of CcO (unpublished results) and electron transport activity; proton pumping has not been estimated) and structural stability (pressure effect on subunit dissociation). We found that high hydrostatic pressure affects the structural and functional stability of monomeric and dimeric CcO quite differently.

The chemical effects on the structure and enzymatic activity of dimeric CcO were investigated (9,13,14). Treat-
ment of dimeric CcO with chemical factors (removal of cardiolipin -14, treatment with urea -13, hydrogen peroxide -9) has important consequences on the structure and function of dimeric enzyme. From our results and considering findings in (14) we can suppose partial dissociation of subunits VIa and VIb due to hydrostatic pressure, which is finished on chromatographic column. In accordance with (14), the decrease in electron transport activity of dimeric CcO due to high pressure was not significant. The exposure of dimeric CcO to chemical factors has more consequences than the exposure to hydrostatic pressure. Possible explanations can be different acting of the pressure. In summary, dimeric form of enzyme is very stable when exposed to high hydrostatic pressure.

The different situation is when the monomeric CcO is treated with high hydrostatic pressure. The exposure of monomeric enzyme to hydrostatic pressure causes dissociation of subunits III, VIa, VIb, VIIa with a significant decrease in electron transport activity. Also the percent of reduction is higher when monomer is exposed to pressure. The subunits VIa and VIb are not responsible for the loss of activity (14) of enzyme. It is known that the dissociation of core subunit III does not alter electron transport activity of CcO (3). From our results we can suppose that the dissociation of subunit VIIa causes significant decrease in enzymatic activity of monomeric CcO. These findings are consistent with results obtained by exposure of CcO to urea (13).

In summary, we can conclude that high hydrostatic pressure perturbs functional (electron transport activity) and structural integrity of monomeric CcO, while dimeric form is more resistant to pressure. Monomeric form of CcO is more fragile than dimeric CcO- this fact can contribute to the general assumption that dimeric form of CcO occurs in vivo.

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