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NO synthesis, unlike respiration, influences intracellular oxygen tension.

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ABSTRACT

We have developed a new phosphorescent probe, PdTCPPNa₄, whose luminescence properties are affected by local variations of intracellular oxygen tension (PO₂). Spectrofluorimetric measurements on living human umbilical venous endothelial cells loaded with this molecule show that a decrease in extracellular oxygen tension induces a fall of PO₂, illustrating the phenomenon of oxygen diffusion and validating the use of this probe in living cells. Moreover, KCN- or 2,4-dinitrophenol-induced modifications of respiration do not lead to detectable PO₂ variations, probably because O₂ diffusion is sufficient to allow oxygen supply. On the contrary, activation by acetylcholine of endothelial nitric oxide synthase (eNOS), that produces NO while consuming oxygen, induces a significant decrease in PO₂ whose amplitude is dependent on the acetylcholine dose, i.e. the eNOS activity level. Hence, activated cytosolic enzymes could consume high levels of oxygen which cannot be supplied by diffusion, leading to PO₂ decrease. Other cell physiology mechanisms leading to PO₂ variations can now be studied in living cells with this probe.

Key-words: endothelium, intracellular oxygen tension, porphyrin, fluorescence
INTRODUCTION

In the cellular respiratory function, part of the energetic metabolism is accompanied by an oxygen intake and a CO₂ discharge in the environment (1). In addition, non-mitochondrial enzymes, not involved in the respiratory function, can also use oxygen: nitric oxide synthase (NOS) is a cytoplasmic enzyme which produces a vasomodulator and neuromodulator gas, nitric oxide (NO), from the precursor L-Arginine, while consuming oxygen (2). In animal cells, extracellular oxygen influx occurs by diffusion through the cell membrane and the intracellular medium, according to the gradient of concentration existing because O₂ is consumed into cells. Using data from the literature (3,4), the maximum oxygen flow entering a cell (diameter: 10µm) by simple diffusion can be calculated in the range of 3×10⁻¹⁵ mol O₂.sec⁻¹.cell⁻¹.

In biological tissues, various means can be used to measure oxygen, such as spectrophotometry (5), oxygen microelectrodes (6,7) and oxygraphy, the latter technique having allowed calculation of basal respiratory O₂ consumption in human umbilical vein endothelial cells (HUVEC): 4×10⁻¹⁷ mol O₂.sec⁻¹.cell⁻¹ (8). Also, non-invasive molecular probes, whose luminescence properties are affected by oxygen, have also been developed to study intracellular (PO₂) or extracellular oxygen tension. For instance, the phosphorescent albumin-bound-Pd meso-tetra (4-carboxyphenyl) porphyrin, has been intravenously injected for in vivo tissue PO₂ measurements in Hamster Skinfold (9) and in rat skeletal muscle (10). Recently, other investigators have measured intracellular PO₂ modifications in response to variations of extracellular PO₂ in isolated dead muscle skeletal fibers (poisoned with 2 mM NaCN) from Xenopus laevis by microinjection of albumin-bound Pd meso-tetra (4-carboxyphenyl) porphyrin (11).

In the present work, we have improved the solubility of the phosphorescent oxygen probe Pd-meso-tetra (4-carboxyphenyl) porphyrin (12) by using a chemically-prepared and purified tetra sodium salt derivative (PdTCPPNa₄). In contrast to the porphyrin microinjection technique used by Lo et al. in their studies on poisoned cells (12), our probe presents the capability of being loaded non-invasively in living cells. On living human umbilical venous endothelial cells (HUVEC), we have studied the relative impacts on PO₂ of two physiological intracellular mechanisms consuming O₂, i. e. mitochondrial respiration and extramitochondrial NO synthesis, together with the effect of hypoxia on
the intracellular PO$_2$, induced by a decrease of the extracellular PO$_2$.

The spectrofluorimetry measurements indicate that both extracellular PO$_2$ decrease and endogenous endothelial nitric oxide synthase (eNOS) activation induce a transient decrease in intracellular PO$_2$, whereas modulation of the mitochondrial respiration does not result in a significant variation of PO$_2$. These findings lead to the hypothesis that maximal O$_2$ diffusion across the cell membrane is sufficient for continuous oxygen supply in mitochondrial respiration, whereas O$_2$ diffusion is insufficient to continuously provide with O$_2$ the activated cytoplasmic enzymes, such as eNOS, that consume high levels of O$_2$. 
MATERIALS AND METHODS

Pd-meso-tetra (4-carboxyphenyl)porphyrin tetra sodium salt derivative.

0.185g of Pd-meso-tetra (4-carboxyphenyl)porphyrin (Porphyrin Products) was neutralized by addition of 13ml NaOH (0.065N). The resulting red solution was stirred for 2 hours at room temperature and filtered, before being evaporated near to dryness (∼0.5ml). The precipitate, Pd-meso-tetra (4-carboxyphenyl)porphyrin tetra sodium salt (PdTCPPNa4), was filtered off, washed with ethanol and dried under vacuum. Yield: 90 %, 169mg.

Intrinsic luminescent properties of the initial molecular phosphorescent probe are not affected by this treatment: emission spectra of PdTCPPNa4 in water solution, obtained with a 514nm laser beam, also shows a maximum emission peak at 710nm (Fig. 1A).

Similar to previously published investigations (13,14), the O2-dependence of the probe phosphorescence is described by the Stern-Volmer relationship: \( \frac{\tau_0}{\tau} = 1 + k_q \tau_0 P_{O_2} \) where \( \tau_0 \) and \( \tau \) are the phosphorescence lifetimes (decay) in the absence of O2 and at a given PO2, respectively, and \( k_q \) (the quenching constant) is a second order rate constant that is related to the frequency of collisions between O2 and the excited triplet state of PdTCPPNa4. An intracellular oxygen tension (PO2) decrease in the environment increases the phosphorescence intensity (I) and lengthens the phosphorescence decay (\( \tau \)) of PdTCPPNa4, and inversely. Here, I and \( \tau \) were followed, as markers of the PO2 variations, in acellular solutions or in PdTCPPNa4–loaded endothelial cells.

Human Umbilical Venous Endothelial Cells (HUVEC).

HUVEC were used for several reasons. The endothelial cells have a glycolytic potential enabling them to stand hypoxia or short duration anoxia (15). Also, endothelial cells allow the investigation of three different mechanisms for inducing potential changes in PO2: i) decrease in extracellular oxygen (by argon bubbling), ii) use of uncouplers and inhibitors of the mitochondrial respiration, and iii) stimulation or blockade of eNOS.

HUVEC were obtained according to a protocole derived from the Jaffe's method and, for investigations of PO2 by fluorescence confocal microscopy, HUVEC were cultured on dishes whose bottom was a glass slide coated with 0.25mg.ml\(^{-1}\) fibronectin (16,17).
Determination of optimal active concentrations of KCN and DNP using oxygraphy technique.

Respiration rate of HUVEC suspended in PSS (2ml, $5\times10^6$ cells.mL$^{-1}$) was measured polarographically in a stirred oxygraph vessel (37°C) including a Clark oxygen electrode. Different concentrations of the respiration inhibitor potassium cyanide (KCN) or of the uncoupler of mitochondrial phosphorylative oxidation 2,4-dinitrophenol (DNP) were then added and the effect on cellular O$_2$ consumption was recorded.

Control of probe penetration in the cells.

Fluorescence images of adherent HUVEC were acquired by using a Zeiss LSM 410 confocal laser scanning microscope (Zeiss). Excitation: 514nm, dichroic 560nm, emission collection: 665nm long pass. Each image (optical sections passed through the nuclei) is the average of 4 successive image acquisition.

Intracellular Oxygen measurement in suspended HUVEC.

Sub-confluent adhering HUVEC were washed twice with PSS and trypsinized. Trypsinization was stopped by addition of PSS containing 50% human serum. Cells were then centrifuged 5min at 200×g and resuspended in 6ml PSS with 0.45mg.ml$^{-1}$ pluronic F-127, 5mg.ml$^{-1}$ BSA added. Sixty microliters of a 10mM PdTCPPNa$_4$ solution were added to a 6ml cell suspension (final concentrations: 100µM PdTCPPNa$_4$) before a 60min incubation at 37°C, 5% CO$_2$, humid atmosphere, in the dark. The cells were then washed of the extracellular fluorescent probe by four centrifugation cycles (5min, 200×g) and pellet resuspension in 6ml PSS containing 4mM octanoate ($\approx10^6$ cells·ml$^{-1}$).

eNOS is activated using 0.1, 1 and 10µM acetylcholine (Ach) (18) or inhibited using 10µM N-ω-nitro-L-arginine methyl ester (L-NAME), an analogue of the precursor L-Arginine which cannot be used by eNOS (19). L-NAME was added to the HUVEC suspension during PdTCPPNa$_4$ incubation (60min). The cell suspension was then washed out of the probe and L-NAME by centrifugation and resuspension in PSS before measurement.

In addition, to evaluate the interaction between PdTCPPNa$_4$ and the chemicals used in this study, the chemicals were added to a 10µM PdTCPPNa$_4$ solution (without cells) during the phosphorescence
emission acquisition.

Two different parameters were measured, depending on the experimental condition: phosphorescence intensity (I) or phosphorescence decay (τ). τ and I were obtained using a pulsed water cooled continuous wave ArGa laser excitation (514nm) and a continuous laser beam (514nm), respectively, and the 90° phosphorescence emission was detected at 710nm with a AsGa photomultiplier.

**Probe and Chemicals.** Chemicals were obtained from Sigma except potassium cyanide (KCN) and 2,4 dinitrophenol (DNP) provided by Merck. PdTCPPNa₄ is prepared as reported above.

**Computer Analysis.** Exponential decay curve analysis was performed by using the Origin 5.0 Microcal Origin Software. For suspended HUVEC intracellular origin measurements, phosphorescence acquisition was performed by using software written by the authors. The averaging of 100 successive exponential decays was used to measure τ.
RESULTS

Optimization of experimental conditions

Validation of the loading protocol

Figure 1B illustrates that PdTCPNa₄ enters human umbilical venous endothelial cells (HUVEC), as seen from the fluorescence confocal microscopy signal (optical slices passing through the nuclei). The weakness of the intracellular fluorescence is expected because of the low quantum yield of this probe. The strong photobleaching due to the repeated laser scanning of the whole microscopic field did not allow further sequential acquisition of images using the confocal microscopy technique. Consequently, all further experiments were carried out on HUVEC suspensions.

Optimal concentrations of KCN and 2.4-dinitrophenol for modulation of mitochondrial respiration in endothelial cells

On HUVEC suspension, using the oxygraphy technique, uncoupling by 100µM DNP stimulates the respiratory rate up to 2-fold within a few seconds, and 1mM KCN is an optimal concentration which induces almost an immediate 10-fold inhibition of respiration. However, 1mM KCN also induces a rapid morphological deterioration of cells observed in confocal microscopy. Since 100µM KCN does not induce cell morphological changes while still inhibiting the respiratory rate by 2-fold (data not shown), 100µM KCN and 100µM DNP have been used as respiration modulators in the following experiments.

Lack of interaction between the PdTCPNa₄ and the chemicals added

It has been verified that acetylcholine and L-NAME do not directly affect the optical properties of PdTCPNa₄ by direct molecular interaction in an acellular physiological solution (Fig. 2A and 2B). Besides, we have observed that the average values of the measured probe phosphorescence decays are different in physiological solution and in cells (about 6μsec and 90μsec, respectively). This can result from the different molecular environment around the probe. In accordance with the literature (12), we have observed, using bovine serum albumin (BSA) addition to PdTCPNa₄ solutions, that the presence of protein induces an increase in the probe luminescence. The same phenomenon likely
occurs when the probe is loaded in the protein-rich cytoplasm of the cell.

Experimental results

Intracellular \( \text{PO}_2 \) variations during modification of extracellular oxygen tension

Argon bubbling (50 min) of a suspended HUVEC loaded with PdTCPPNa\(_4\) was used to deplete the extracellular medium of solubilized \( \text{O}_2 \), and phosphorescence intensity was recorded. PdTCPPNa\(_4\) phosphorescence intensity increased by 7 fold, plateauing after 25 min (Fig. 3). This indicates that PdTCPPNa\(_4\) strongly responds to the intracellular \( \text{PO}_2 \) decrease induced by extracellular decrease in oxygen tension.

Mitochondrial respiration and \( \text{PO}_2 \) variations

Under these conditions, the measurement of phosphorescence decay (\( \tau \)) avoids the problem of working with the very weak signal of PdTCPPNa\(_4\). As shown in figure 4, the addition of 100\( \mu \)M DNP or 100\( \mu \)M KCN does not modify the initial value of the phosphorescence decay (\( \tau \)) over 30 min in PdTCPPNa\(_4\)-loaded HUVEC, suggesting that mitochondrial respiration does not significantly influence \( \text{PO}_2 \).

Non-respiratory oxygen consumption by eNOS and variations of \( \text{PO}_2 \)

Addition of 0.1, 1 and 10\( \mu \)M acetylcholine to HUVEC induce transient and dose-dependent increases in \( \tau \) (Fig. 5A, Fig. 6A, 6B, 6C), generally occurring after 2-5 minutes (up to 10 min in a few experiments), whereas no increase in \( \tau \) is observed in the corresponding control experiments (Fig. 5B). These results suggest a significant reduction of the intracellular \( \text{PO}_2 \) under acetylcholine stimulation. The average percentages of increase in amplitude of the phosphorescence decay peaks, as compared to baseline, are in the range of 4.2\% (0.1\( \mu \)M Ach), 23\% (1\( \mu \)M Ach) and 81.6\% (10\( \mu \)M Ach) (Fig. 6D). A statistical analysis using a one-way ANOVA showed that the average percentages of increase in the amplitude of the phosphorescence decay peaks are significantly dependent upon the acetylcholine dose (\( P<0.05 \)).
Figure 5C shows that there is no increase in \( \tau \) in response to 1\( \mu \)M acetylcholine immediately after incubation with L-NAME. This tracing is comparable to the control (Fig. 5B). Nevertheless, 2 hours after the end of L-NAME incubation, after eNOS activity resumed, the acetylcholine-induced \( \tau \) elevation –therefore PO\(_2\) decrease- is restored (Fig. 5D). The fact that no detectable modification of luminescence, following Ach addition on L-NAME-treated HUVEC, indicates a causal relationship between O\(_2\) consumption related to eNOS activation and the transient increase in \( \tau \). Moreover, the literature mentioned that a few porphyrins are sensitive to NO (20) or can be used as NO probe (21). Consequently, 10 and 100nmol NO (final concentrations: 3\( \mu \)M and 30\( \mu \)M, respectively) have successively been added to a PdTCPNa\(_4\) solution (10 \( \mu \)M) in PSS, in order to verify that NO does not modify the value of \( \tau \) corresponding to our probe (Fig. 2C). Therefore, these results depict actual PO\(_2\) modifications rather than NO level detection.
DISCUSSION

The present in vitro study validate a novel technique for measuring intracellular PO\textsubscript{2} variations in acellular solutions as well as in living cells. Although, values of basal phosphorescent decay in acellular solutions (Fig. 2) and in living cells (Figs. 4 and 5) are different because the local environment surrounding the probe in the cell is likely to affect its response to PO\textsubscript{2} variations. PdTCPPNa\textsubscript{4} loaded into human umbilical venous endothelial cells (HUVEC) exhibits spectral properties similar to the well characterized phosphorescent oxygen probe Pd meso-tetra (4-carboxyphenyl) porphyrin (12). Moreover PdTCPPNa\textsubscript{4} could be used for the measurement of intracellular oxygen tension (PO\textsubscript{2}) variations in living HUVEC, as shown by the intracellular response to extracellular PO\textsubscript{2} decrease induced by argon bubbling (Fig. 3). The original loading protocol used in this study, in contrast with microinjection techniques which allows measurements in a few cells at a given time (11), enables the study of intracellular PO\textsubscript{2} in a population of millions of cells. Furthermore, microscopy studies do not show any morphological cell damage due to PdTCPPNa\textsubscript{4} loaded in HUVEC, even under focused light (Fig. 1B). In addition, the fact that low intracellular concentrations of PdTCPPNa\textsubscript{4} were used is likely to keep the effect of any reactive O\textsubscript{2} species produced, due to energy transfer from the porphyrin probe in its excited state to molecular oxygen, at a very low level.

Extracellular oxygen transport towards the inside of the cell occurs by diffusion according to the gradient of concentration and tends to maintain a constant PO\textsubscript{2}. Using data from the literature (3,4), theoretical maximum oxygen flow entering the cell by diffusion can be estimated in the range of 3\times10^{-15} mol O\textsubscript{2}. sec\textsuperscript{-1}.cell\textsuperscript{-1} and oxygen uptake measured by oxygraphy in HUVEC (37°C) has been found at a basal level, about 75 times lower: 4\times10^{-17} mol O\textsubscript{2}.sec\textsuperscript{-1}.cell\textsuperscript{-1}(8). Therefore, physiological modulations of mitochondrial respiration, which usually vary by a factor 2-4 times (8), should have no impact on intracellular PO\textsubscript{2}, because O\textsubscript{2} diffusion flow allows a O\textsubscript{2} influx 20-50 fold higher. Thus, it is not surprising that, in these conditions, modulation of mitochondrial respiration using 100µM KCN or 100µM DNP, which respectively inhibits and stimulates respiratory function by 2-fold, does not measurably modify PO\textsubscript{2} (Fig. 4).

Besides, oxygen consumption independent from respiration and resulting from activation of endothelial nitric oxide synthase (eNOS) has been evaluated. The use of a NO sensitive
microelectrode previously allowed to calculate NO concentration, at a distance of 50µm from the surface of rat aorta endothelial cell, in the range of $6.2\times10^{-16} \text{ mol NO}\cdot\text{sec}^{-1}\cdot\text{cell}^{-1}$ (22). Therefore, since NO synthesis has a stoichiometry of 1/1 with regard to consumed O$_2$, the basal production of NO by eNOS in rat aorta single endothelial cell requires the consumption of about $6.2\times10^{-16} \text{ mol O}_2\cdot\text{sec}^{-1}\cdot\text{cell}^{-1}$. Moreover, eNOS has been shown to be activated by endothelial cell stimulation with acetylcholine (Ach), leading to activation of NO release (23). In the endothelium from rabbit aorta, Ach increases NO extracellular level -therefore production- by $\approx 10$ times (23) and probably more if intracellular NO degradation is taken into account. Therefore, the corresponding quantity of O$_2$ transiently consumed by Ach-stimulated eNOS can be evaluated in the range of 10 times the basal consumption, i.e. $6.2\times10^{-15} \text{ mol O}_2\cdot\text{sec}^{-1}\cdot\text{cell}^{-1}$. This value is about 2 fold higher than the maximum rate of O$_2$ diffusion through the cell membrane. Consequently, eNOS activation is a physiological mechanism which can induce a much stronger decrease in intracellular PO$_2$ than modulation of respiration. Our experiments corroborate this calculation, since 0.1-10µM acetylcholine induce transient and dose-dependent increases in PdTCPPNa$_4$-loaded cell phosphorescence decay, indicating a corresponding PO$_2$ reduction (Fig. 5A and Fig. 6). On the contrary, no change in porphyrin phosphorescence was observed when cell respiration was increased or decreased by the use of DNP or KCN (Fig. 4). Also, as expected, no PO$_2$ variation is detected in response to acetylcholine when eNOS is blocked by L-NAME (Fig. 5 C and D).

The present results suggest that other oxygen-related mechanisms of the cell physiology can now be further investigated using cell loading with PdTCPPNa$_4$. 
REFERENCES


FIGURE LEGENDS

Figure 1. Emission spectrum of the water soluble PdTCPPNa₄ illuminated by a 514 nm laser beam (A.). Adherent HUVEC loaded with PdTCPPNa₄ observed by confocal laser scanning microscopy in differential interferential contrast (left) and fluorescence (right) (B.). The images presented are representative of two experiments.
Figure 2. Absence of interaction of NO, Ach and L-NAME on the luminescence properties of PdTCPPNa₄ (10µM) in PSS. (A.) Effects of 1µM acetylcholine or 10µM L-NAME on the 10µM porphyrin solution. (B.) Effects of 1µM acetylcholine and 10µM L-NAME on the 10µM porphyrin solution and corresponding control tracing. (C.) Sensitivity of the porphyrin probe (10µM) to successive additions of 10 and 100nmol NO. Arrows: product addition. A representative tracing is shown (n=2 in each case).
Figure 3. Effect of argon bubbling on the phosphorescence intensity of PdTCPPNa₄-loaded HUVEC suspension. The argon bubbling starts at time 0min.
Figure 4. Effect of 100µM DNP (A.) and 100µM KCN (B.) on the phosphorescence decay of 100µM PdTCPPNa$_4$-loaded HUVEC suspension. A control tracing is shown in C. A representative tracing is shown (n=2 in each case).
Figure 5. 1µM acetylcholine (Ach) induces a transient lengthening of phosphorescence decay (τ) on 100µM PdTCPNa₄-loaded HUVEC (A.). Control tracing (B.). Effect, on 100µM PdTCPNa₄-loaded HUVEC, of 1µM Ach immediately after incubation with 10µM L-NAME (C.) and 2 hours after the end of L-NAME incubation (D.). A representative tracing is shown in each case (A: n=3 ; B,C,D: n=2).
Figure 6. Dose-effect of acetylcholine (Ach) on the phosphorescence decay ($\tau$) of 100µM PdTCPPNa$_4$-loaded HUVEC. The amplitude of the phosphorescence decay peak induced by Ach addition increases with Ach concentration: 0.1 µM (A), 1 µM (B) and 10 µM (C). A representative tracing is shown in each case (A: n=5 ; B and C: n=4). The arrows indicate the time of Ach addition (A, B and C). The average percentages of increase in amplitude of the phosphorescence decay peaks induced by Ach, as compared to baseline, were calculated for each Ach dose using the results from all experiments, and represented in D.