Diphenyleneiodonium, an inhibitor of NOXes and DUOXes, is also an iodide-specific transporter

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ABSTRACT

NADPH oxidases (NOXes) and dual oxidases (DUOXes) generate O2– and H2O2. Diphenyleneiodonium (DPI) inhibits the activity of these enzymes and is often used as a specific inhibitor. It is shown here that DPI, at concentrations similar to those which inhibit the generation of O2 derivatives, activated the efflux of radioiodide but not of its analog 99mTcO4– nor of the K+ cation mimic 86Rb+ in thyroid cells, in the PCC3 rat thyroid cell line and in COS cell lines expressing the iodide transporter NIS. Effects obtained with DPI, especially in thyroid cells, should therefore be interpreted with caution.

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1. Introduction

Seven specific O2– and H2O2 generating enzymes have been demonstrated, the NADPH oxidases, NOXes and dual oxidases DUOXes. In many cell types, they generate both intracellular and extracellular O2– and H2O2. They share many properties, including enzymatic mechanisms, but differ in terms of function, tissue distribution and control [1–3].

Diphenyleneiodonium (DPI) inhibits the activity of these flavoenzymes. It is often considered as a specific inhibitor. Inhibition of any metabolic step by DPI is therefore often attributed to this effect [4–9]. Although caution has been advised against the use of inhibition by DPI as a specific biomarker of NOX and DUOX actions [4], such arguments are still commonly used [10].

In this article we demonstrate another action of DPI which raises new interesting questions. In the course of our study of iodide metabolism in thyroid cells we observed that it increased the release of 125I iodide but not 99mTcO4–, thus behaving as a specific carrier of iodide.

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This shows an effect of DPI other than the inhibition of H2O2 generation. It presents an interesting chemical property of this iodinated derivative and may suggest a therapeutic application.

2. Materials and methods

2.1. Reagents

125I as NaI (~17Ci/(629GBq)/mg) and 86Rb+ (19.3 mCi/(714.1 MBq)/mg) were purchased from Perkin Elmer Life and Analytical Science (Zaventem, Belgium). 99mTc as Na Pertechnetate (99mTcO4–) was provided to us by the Medical School Hospital. It was prepared daily, carrier free, from a 99mTc generator (Ultra TechneKow, Mallinckrodt Nuclear Medicine). We used around 1 μCi tracer per dish.

Homovanillic acid (HVA), horse radish peroxidase type II (HRP), methylmercaptomidazole (MML) and diphenyleneiodonium chloride (DPI) were provided by Sigma–Aldrich BVBA (Diegem, Belgium). Sodium iodide and sodium perchlorate were purchased from Merck KGaA (Darmstadt, Germany).

2.2. Tissue

Pig thyroids were collected at a local slaughterhouse. The fresh tissue was sliced at room temperature with a Stadie-Riggs microtome (Arthur Thomas, Philadelphia, PA, USA). Slices (0.3 mm thick, 45–60 mg wet weight) were incubated in 2 ml incubation medium.
2.3. Cell culture

PCCl3 cells, a rat thyroid cell line, were grown in Coon's modified Ham's F12 medium supplemented with 5% decomplemented fetal bovine serum, bovine TSH (1 μU/ml), transferrin (5 μg/ml), insulin (1 μg/ml), penicillin (100 IU/ml), streptomycin (100 μg/ml) and fungizone (2.5 μg/ml). COS–NIS cells are COS-7 cells transfected with the wild type human sodium/iodide symporter (NIS) cDNA. Stable cell lines were selected by resistance to geneticin [11]. They were cultured in DMEM supplemented with 10% fetal bovine serum, Na pyruvate (1 mM), penicillin (100 IU/ml), streptomycin (100 μg/ml) and fungizone (2.5 μg/ml).

2.4. Incubation

The slices and cells were incubated in Krebs–Ringer Hepes medium (NaCl 124 mM; KCl 5 mM; MgSO4 1.25 mM; CaCl2 1.45 mM; Hepes Buffer 25 mM) supplemented with glucose 8 mM and BSA (bovine serum albumin) 0.5 mg/ml [12].

2.5. Measurement of the H2O2 production

Thyroid slices were preincubated 1 h in KRH medium and then transferred to fresh medium containing 0.1 mg/ml HRP, 440 μM HVA and the tested agonists for 90 min. When H2O2 is produced by the tissue, the HVA is oxidized into a fluorescent dimer by the peroxidase. The fluorescence of the medium was measured at 315 and 425 nm, excitation and emission wavelengths respectively [13]. Generation of H2O2 was expressed as ng H2O2 produced by 100 mg of tissue, or ng H2O2 per dish in the case of cells, during the length of the incubation.

2.6. Measurement of iodide uptake

The thyroid slices were preincubated 1 h in KRH supplemented with 0.1 mM MMI to block organization of iodide, then incubated for 1 h with fresh KRH medium supplemented with 125I− (1 μCi/ml), 10−7 M KI, 0.1 mM MMI and the tested agents, washed and counted. The uptake was evaluated by the ratio: counts per min 125I−/100 mg tissue to counts per min 125I−/100 μl medium [14]. Uptake of investigated cells in culture was divided by the uptake of cells incubated in the presence of NaClO4 1 mM to obtain concentration ratios (cell to medium C/M).

2.7. Measurement of tracer efflux from cells

300,000 cells were seeded in 35-mm-diameter dishes and grown 24 h in the case of COS–NIS cells or 48 h in the case of PCCl3 cells, in their respective culture medium. The cells were then rinsed and incubated under slight agitation in KRH medium supplemented with KI 10−7 M and 1 μCi/ml 125I−, or 1 μCi/ml 99mTcO4−, or 1 μCi/ml 86Rb+. As no iodination takes place in COS or PCCl3 cells, MMI was not used routinely in these experiments. For 86Rb+ uptake, the K+ concentration in the medium was 0.5 mM instead of 5 mM.

After 45 min uptake with 125I− or 99mTcO4− and 2 h with 86Rb+, the cells were rinsed twice with KRH and incubated with fresh medium, which was supplemented with KI (10−7 M) and ClO4− (1 mM) in the case of 125I− or 99mTcO4− efflux, to avoid tracer recirculation. DPI (10 μM) is added at the start of this second incubation. Fifty microliter from 1 ml medium was withdrawn at the mentioned times and counted. At the end of incubation, the cells were rinsed twice with KRH medium kept on ice, then dissolved in 1 M NaOH and counted. The efflux was expressed in percent of the total radioactivity taken up by the cells (cells + medium + serial aliquots).

3. Results

DPI, in the same concentration range, inhibits both H2O2 generation and radioiodide uptake in pig thyroid slices (Fig. 1).

The effect of DPI on iodide uptake level at equilibrium results from the balance between the continuous uptake by NIS and the release. The inhibition of iodide uptake by DPI takes place at a lower concentration (IC50 1–4 μM) than the effect of competing iodide (IC50 40 μM) (Fig. 2). Moreover the C–I bond of the iodine to the benzene ring of DPI is not labile [9]. The inhibition therefore is not due to iodide release from DPI.

For the release experiments (Fig. 3, 4, 5), DPI is added after the labeling of the cells and the release is measured immediately after; it is, in effect, quasi instantaneous. The effect of DPI does not correspond to a general membrane permeabilization as preloaded 86Rb+, used as analog of K+, is not discharged from PCCl3 nor COS–NIS cells (Fig. 3).

Moreover, DPI is a carrier more specific than NIS, the transporter
Fig. 3. Comparison of DPI effect on $^{125}$I$^-$ and $^{86}$Rb$^+$ efflux from COS–NIS preloaded cells. The efflux of the tracer from the cell to the medium is expressed as percentage of total uptake by the cells. The $^{125}$I$^-$ uptake at equilibrium was 24.6 in this experiment. (Ratio of control vs NaClO$_4$ treated cells). Empty circles: control. Filled square: DPI 10 μM.

Fig. 4. Comparison of DPI effect on $^{125}$I$^-$ and $^{99m}$TcO$_4^-$ efflux from PCCl$_3$ and COS–NIS preloaded cells. DPI was added at the beginning of the efflux. The uptake values at equilibrium for $^{125}$I$^-$ were 24 ± 1 and 24 ± 0.05 in PCCl$_3$ and COS–NIS cells respectively. The uptake values at equilibrium for $^{99m}$TcO$_4^-$ were 216 ± 6 and 228 ± 9 in PCCl$_3$ and COS–NIS cells respectively (mean of duplicates). The results of $^{125}$I$^-$ and $^{99m}$TcO$_4^-$ efflux are expressed as percentage of total uptake by the cells. Empty circles: control. Filled square: DPI 10 μM.
of iodide (Na⁺/I⁻ symporter) as it does not release ⁹⁹ᵐTcO₄⁻ from prelabeled cells. Thus, contrary to the NIS [15], DPI does not recognize ⁹⁹ᵐTcO₄⁻ as radiiodide (Fig. 4).

Spontaneous iodide release is much decreased when the temperature is lowered from 37 to 22 °C and 15 °C (Fig. 5). However, the relative effect of DPI is as important at the three temperatures.

To check the reversible or permanent character of DPI effect on iodide uptake (through its effect on efflux) and on H₂O₂ production we designed a protocol made of 3 sequential incubations called A, B and C. The measurement of iodide uptake (in the presence of ¹²⁵I and 10⁻⁷ M KI, for 45 min) and of H₂O₂ production (in the presence of 440 μM HVA and 100 μg/ml Peroxidase II for 90 min), took place in the last incubation C. Incubation A and B lasted 45 min each for either parameter. No DPI was added in any of these incubations, for the control. To test if DPI effect was short or long lasting we added it at 3 different times: In A only, or in B only, or in C only. The results presented in Fig. 6 show that the effect of DPI on iodide uptake is strongest when added in C, but fade away if it is added in B only and more so if added in A only, which suggests that the DPI involved in iodide efflux is rapidly washed in the absence of DPI.

The effect of DPI on H₂O₂ is much more persistent. Indeed, if added previously, even in the first incubation A only, it works as well. This suggests a more stable binding of the DPI involved in this inhibition.

4. Discussion

We have shown that DPI, at concentrations of the same order as those that inhibit H₂O₂ generation by thyroid cells, also causes an immediate specific release of iodide from thyroid and non-thyroid cells.

The effect of NaClO₄ or NaSCN which only compete with the uptake of iodide and decrease by this mechanism the cell/medium equilibrium level [15] is relatively slow. The immediate effect of DPI on iodide efflux carried out in the presence of perchlorate is therefore not due to an inhibitory role on iodide uptake by NIS. It does not seem to require the specific iodide channel(s) involved in the efflux of iodide at the apical membrane of the thyrocyte [16,17], as it is reproduced in COS–NIS cells i.e. in non-thyroid cells.

Besides, an effect on NIS would similarly affect the efflux of ⁶⁰ᵐTeO₄⁻ [15,18]. The absence of effect on ⁶⁰ᵐTeO₄⁻ or ⁸⁶Rb⁺ release, excludes a general permeabilization of the cells by DPI.

Other unspecified effects of DPI have been observed but at higher concentrations than those effective on NOXes. For example, DPI (10 μM) also directly inhibits cholinesterase activity and to a lesser extent an internal Ca⁺⁺ pump [19]. The effect of DPI on iodide efflux is not due to the inhibition of H₂O₂ generation as it occurs in COS–NIS cells without DUOX and is, contrary to the effect on DUOX, relieved by incubation without the drug.

We have no explanation for this effect on iodide. Certainly because the covalent link in DPI between iodine and the other part of the DPI molecule is not labile, there is no dynamic exchange with medium iodide. The molecule is stable under stringent conditions [9].

Moreover, higher concentrations of iodide than DPI are needed to reach the same inhibition of C/M.

We can therefore only speculate on the mechanism of the DPI effect. A first explanation would be the rapid activation of an anion channel in the cell [20]. However DPI also acts on COS–NIS cells [11] which do not express any iodide specific channel, especially a channel distinguishing iodide and ⁹⁹ᵐTcO₄⁻. A second explanation could be that DPI behaves as a membrane ionophore specific for iodide.

Catalysis of an anion/hydroxyl ion exchange across mitochondrion membrane by DPI has been reported [21]. The fact that depressing membranes fluidity by decreasing the temperature of the cells slow down both basal and the increased efflux, does not distinguish the hypothesis of DPI acting as a shuttle for iodide from the concept that it acts as an intramembrane channel as is the case for valinomycin
and K⁺ [22,23]. An ionic type of carrier mechanism would be similar to the mitochondrial protein carrier mechanism which is believed to account for the uncoupling effect on mitochondrial oxidative phosphorylation by dinitrophenol. Both hypotheses of channel and shuttle, would suggest a specific affinity of DPI molecule for iodide.

Nevertheless the present data confirm that DPI cannot be considered as a specific inhibitor, of NOXes and DUOXes. They suggest that DPI or an analog with similar properties could therefore be considered as drugs for iodine or amiodarone induced hyperthyroidism as they would combine in one molecule the inhibition of thyroid hormone synthesis and iodide uptake (in fact retention) as a combination of antithyroidperoxidase drugs and perchlorate.

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