Pulmonary Oxidative Status in Norma and Pathologies on the Basis of Analysis of Exhaled Breath Condensate

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Abstract

On the basis of analysis of the exhaled breath condensate (EBC), which is truly non-invasively collected, we propose to evaluate the pulmonary oxidative status. As known, both peroxides, primarily hydrogen peroxide (H₂O₂), and nitrite ions (NO₂⁻) present in EBC. However, where as peroxides serve as strongest oxidants, nitrite in the same medium due to its electrochemical potential can only be a reductant. We have shown, that H₂O₂ concentration in EBC being at micromolar level for healthy human is increased in case of inflammatory pathologies (asthma, pneumonia, bronchiectases). Moreover, monitoring H₂O₂ in EBC one can evaluate the degree of inflammation (exacerbation vs remission) and even improve the personified therapy. On the contrary, nitrite concentration in case of similar inflammatory deceases is decreased. A simultaneous increase of oxidant (H₂O₂) and a decrease of reductant (NO₂⁻) concentrations in EBC point to a significant pulmonary oxidative stress accompanying inflammatory pathologies.

Keywords: Exhaled breath condensate; hydrogen peroxide; nitrite; inflammation; oxidative stress.

1. Introduction

Non-invasive diagnostics attracts nowadays a growing interest. However, there are only a few approaches allowing truly non-invasive quantitative evaluation of key blood metabolites. Among them the minimal traumatic for patients provides the collection of exhaled breath condensate (EBC). Appearance in clinical practice of commercial EBC collectors has led to standardization of this method.

Among the great advantages of EBC is a possibility to deliver a kind of portrait of metabolites. Indeed, due to a minor presence of proteins, the biocatalytic activity in EBC is
negligibly low. Hence, even the ratio of redox metabolites reminds it in vivo.

Hydrogen peroxide (H₂O₂) is known to be the strongest oxidant in organisms, characterized in physiological solutions by the highest oxidative potential, which is even higher than that of superoxide radical (O₂⁻) [1]. In contrast to the superoxide, H₂O₂ is stable in aqueous solutions, and hence can be considered as a valuable marker for oxidative stress.

Existence of hydrogen peroxide in exhaled breath condensate has already been reported by a number of authors [2-4]. The main problem with the existent data, however, is a remarkable variation of H₂O₂ concentrations reported even for healthy human. The lowest reported values are less than hundreds of nano-moles per liter [5], whereas some authors claim several micro moles per liter content of hydrogen peroxide in EBC [6, 7]. As seen, the amplitude is almost two orders of magnitude.

Another problem with the reports on hydrogen peroxide concentration in EBC is the generally accepted it’s increase in case of nearly all lung diseases [8]. However, where as for inflammatory diseases (asthma, pneumonia etc.) the increase of H₂O₂ in EBC [9-11] is expected due to oxidative radical formation as organism response, the pathologies like chronic obstructive pulmonary disease (COPD) are not directly accomplished with the free radical formation mechanism. Nevertheless, even for COPD the remarkable raise in hydrogen peroxide concentration is reported [12, 13].

We already reported on Prussian Blue as the most advantageous hydrogen peroxide transducer [14-16]. Comparing with the most widely used platinum, Prussian Blue modified electrodes are: (i) three orders of magnitude more active in H₂O₂ reduction and oxidation in neutral media, and (ii) three orders of magnitude more selective for hydrogen peroxide reduction in the presence of oxygen [17]. The attractive performance characteristics of the electrochemically deposited Prussian Blue allowed to denote it as artificial enzyme peroxidase [15, 16]. Moreover, on the basis of nano-structured Prussian Blue (its nano-electrode arrays) the electrochemical sensor with the record performance characteristics has been elaborated [18, 19]. Hence, it was highly attractive to apply the advanced sensor for hydrogen peroxide to monitor H₂O₂ in EBC.

Except for H₂O₂, a number of metabolites were found in EBC. Our interest attracted nitrite (NO₂⁻), an ion closely chemically related to nitric oxide (NO), which is known to have a great biological importance [20]. Nitrite can serve as both oxidant and reductant and, hence, can also be considered as a marker for oxidative stress. Nitrite found to be at a level of 0.33-3.3 micromoles per liter of EBC in normal non-smokers [21, 22]. In literature the concentration of nitrite in EBC is increased in case of any pulmonary disease [23-25].

We, however, note that a potential of (NO₃⁻/NO₂⁻) redox couple is almost equal to normal hydrogen potential (0.01 V, NHE), which is dramatically lower compared to redox potential of (H₂O₂/H₂O) reaction (1.77 V, NHE) [26]. For comparison, the voltage required for water decomposition into hydrogen and oxygen is just 1.23 V. Hence, nitrite versus H₂O₂ as metabolite of oxidative stress can be considered only as a reductant, rather than an oxidant. This makes doubtful the reported increase of both H₂O₂ and NO₂⁻ in EBC in case of inflammatory diseases.

In the present work we combined efforts of clinicians and analytical chemists to evaluate pulmonary oxidative status on the basis of EBC analysis. We’ve proven that hydrogen peroxide concentration in EBC of healthy human exists at a level of micro-moles per liter. Low-inflammatory pathologies (COPD) cause a small increase of H₂O₂ content in EBC. On the contrary, the main inflammatory diseases (asthma, pneumonia etc.) are accomplished by a significant (up to several times) increase of hydrogen peroxide content in EBC. A simultaneous increase of H₂O₂ and decrease of NO₂⁻ content in EBC points to a significant pulmonary oxidative stress upon inflammatory pathologies.

2. Experimental

2.1 Subjects

Four groups of in-patients (n=151) were included in the present study: COPD patients with stage II-III disease (n=70); patients with moderate-to-severe asthma (n=44); patients with moderate
community-acquired pneumonia (n=24); and patients with bronchiectases (n=13).

The diagnosis of COPD, asthma, pneumonia and bronchiectases were based on the generally accepted (standard) criteria [27-29].

The control group included 74 subjects (age range 24–34 yrs; mean±SD 24±5 yrs), including 27 healthy smokers without symptoms (St George’s questionnaire for symptoms such as cough and sputum production) whose lung function parameters were normal. Atopy and allergies were excluded.

The Ethics Committee of Research Institute of Pulmonology, Moscow, Russia, approved the study protocol, and all subjects gave informed consent before participating.

2.2 Chemicals

Experiments were carried out with MilliQ water from a Millipore MilliQ system. All inorganic salts and hydrogen peroxide (30% solution) were obtained at the highest purity from Reachim (Moscow, Russia) and used as received.

Horse radish peroxidase (HRP), 2,2’-azino-bis(3-ethylbenzthiazoline-6-sulfonic acid) (ABTS), catalase from bovine liver were purchased from Sigma (St Louis, USA). Aldrich (Steinheim, Germany).

Exhaled breath condensate (EBC) samples were collected using a commercially available condenser ECoScreen® (Erich Jaeger GmbH, Germany) in morning hours. After rinsing their mouths, subjects breathed tidally through a mouthpiece and a two-way nonrebreathing valve, in which inspiratory and expiratory air were separated, and saliva trapped. They were asked to breathe at a normal frequency, wearing a nose clip, for a period of 10 min to obtain approximately 1.5 ml condensate. All collected samples were immediately stored at -70°C and kept for less than a week before analysis.

2.3 Hydrogen peroxide detection

Hydrogen peroxide concentration in EBC was detected in flow-injection system equipped with Prussian Blue based sensors. The flow injection system consisted of a Cole Parmer (Vernon Hills, IL) peristaltic pump (7519-10), homemade flow-through wall-jet cell with 0.5-mm nozzle positioned at 1±2-mm distance from the surface of disk electrode, (Ag|AgCl)l M KCl) reference, homemade injector, and PalmSens potentiostat (Palm Instruments BV, Netherlands) interfaced to an IBM PC. Flow rates used were in the range 0.5±1 mL min\(^{-1}\). In FIA experiments, the peak current values were taken for data treatment, sample volume was 50 µl, and working electrode potentials were 0.00±0.05 V, allowing hydrogen peroxide reduction on Prussian Blue-modified electrodes. Carrier solution was 0.05 M phosphate buffer (pH 6.0) containing 0.1 M KCl. EBC prior to injection was diluted 3 times with the carrier.

Prussian Blue based sensors for hydrogen peroxide were elaborated on the basis of screen printed three-electrode chips with carbon both working and counter electrodes, and Ag|AgCl reference. Prussian Blue was deposited in cyclic voltammetric conditions with switching potentials of 0.3±0.4 V (cathodic) and of 0.7±0.8 V (anodic) at a sweep rate of 40 mV s\(^{-1}\), or by applying a constant potential of 0.4 V as described elsewhere [16]. Growing solution contained 4 mM K\(_3\)[Fe(CN)\(_6\)] and 4 mM FeCl\(_3\). A solution of 0.1 M HCl and 0.1 M KCl was used as supporting electrolyte. After deposition, Prussian Blue films were electrochemically activated in the same supporting electrolyte by cycling in the range -0.05±0.35 V at a rate of 40 mV s\(^{-1}\) until a stable voltammogram was obtained. Then the electrodes were heated at 100°C for 1 hour.

Spectrophotometric hydrogen peroxide detection was carried out using LKB-Ultraspec VII (Bromma, Sweden) spectrophotometer. Concentration of peroxidase and ABTS (extinction coefficient (\(\varepsilon\)) of 35 mmol L\(^{-1}\) cm\(^{-1}\) at 405 nm) in phosphate buffer (pH 5.5) was 5·10\(^{-9}\) mol L\(^{-1}\) and 5·10\(^{-4}\) mol L\(^{-1}\), respectively. Calibration was made considering initial rates of peroxidase-catalyzed ABTS oxidation by hydrogen peroxide at certain concentrations of the latter. Peroxidase concentration was independently controlled at 403 nm (\(\varepsilon\)=1.05·105 mol L\(^{-1}\) cm\(^{-1}\)).

2.4 Nitrite detection

The detection principle was the diazotization of the end toluidine groups of polyurethane foam by nitrite ions in acidic medium with subsequent
coupling with 3-hydroxy-7,8-benzo-1,2,3,4-tetrahydroquinoline [30]. The latter was synthesized as follows. 1-Naftilamine (42.9 g, 0.3 mol) and epichlorhydrin (27.9 g, 0.3 mol) in 40 ml of 1,2-dichlorobenzene were shaken at 160 °C for 7 hours. The precipitate was washed with PhCl and hexane subsequently (10 ml of each). Then the product was suspended in 150 ml of 25% aqueous NH₃ and washed with water to neutral pH. The yield of the final product was 60%.

The product of diazo coupling with intensive orange color was monitored at 470 nm with the diffusion reflection spectrometer (“Spectrotron”, Chimautomatika, Russia). Polyurethane foam was purchased from “Radical” (Kiev, Ukraine).

Nitrite detection in EBC was made as follows. One milliliter of EBC was mixed with 1 ml of 5 M HCl and 3 ml of water. Afterwards a tablet of polyurethane foam was immersed in this solution and shook for 30 min. The tablet of diazotized polyurethane foam was dried with blotting paper and then immersed in aqueous solution of 100 µg 3-hydroxy-7,8-benzo-1,2,3,4-tetrahydroquinoline in 5 mL of 0.2 mol L⁻¹ Na₂CO₃. After 30 min of shaking the tablet was washed and the diffusion reflection spectra were recorded. Nitrite concentration in EBC was estimated from calibration graph.

2.5 Statistical analysis

Data are presented as mean±SD for not normally distributed data. Data for all groups were analyzed by the Kruskall-Wallis test and differences between individual variables from two groups were analyzed by the Mann–Whitney U-test. Correlations between variables were sought using the Spearman rank correlation test. A p-value <0.05 was considered significant.

![Figure 1](image_url)

**Figure 1.** Concentration of hydrogen peroxide (H₂O₂) in EBC of healthy people and patients with different pulmonary diseases.
3. Results and discussion

Hydrogen peroxide (H_{2}O_{2}) is already well known as a marker for different pulmonary pathologies. Accordingly, exhaled breath condensate from different groups of pulmonary patients as well as for healthy humans was first analyzed concerning the presence of hydrogen peroxide. Considering huge variation of the reported concentrations of H_{2}O_{2} in EBC (the amplitude is almost two orders of magnitude), we decided first to determine its true level.

As seen in figure 1, the flow-injection (FIA) system equipped with Prussian Blue based hydrogen peroxide sensor responds to EBC as to H_{2}O_{2} solutions of micro-Molar concentrations. As published elsewhere, FIA systems equipped with Prussian Blue based sensors usually display the lower limit of the calibration range below 0.1 µmol/L [15-19]. To confirm, that the response of FIA system was indeed to hydrogen peroxide, the enzyme catalase (which catalyses dismutation of hydrogen peroxide) in concentration of 5 µg per ml was added to EBC. After treatment with catalase the response of FIA system to EBC was decreased by 40-60%. Hence, the main part of the FIA system response is indeed generated by free hydrogen peroxide in EBC. The fact, that treatment with catalase did not cause a complete disappearance of the signal in FIA system, can be explained in terms of complex formation between H_{2}O_{2} and organic compounds (like urea). Alternatively, Prussian Blue based sensors can respond to organic peroxides present in EBC.

The reference method for hydrogen peroxide detection in EBC was the peroxidase enzyme spectrophotometric test. The reference gave similar results to flow-injection system (data not shown). We note that peroxidase enzymes are not specific just to hydrogen peroxide, and catalyze reduction of some organic peroxides as well. Hence, from both electrochemical and spectrophotometric investigations we cannot exclude peroxides presented in EBC in form of organic peroxides. We were unable to find analytical tools to discriminate hydrogen peroxide from organic peroxides in EBC and thus refer to total peroxide content as to hydrogen peroxide concentration.

From this part we conclude, that EBC collected with EcoScreen contains H_{2}O_{2} at micro-Molar level.

We determined hydrogen peroxide in EBC of healthy human and of different groups of pulmonary patients. As seen in fig. 1, hydrogen peroxide concentration in EBC of healthy human is at a level of 1.5 µmol L^{-1}. Smokers have a general tendency for increase of H_{2}O_{2} content in their EBC, but the difference with non-smokers is not valuable.

All pulmonary patients are characterized by increased H_{2}O_{2} concentration in EBC (fig. 1). However, for low-inflammatory disease like COPD the increase of H_{2}O_{2} content is not valuable. On the contrary, inflammatory pathologies (asthma, pneumonia, bronchiectases) cause the significant increase of H_{2}O_{2} concentration in EBC (fig. 1). In certain cases the increase of H_{2}O_{2} content versus healthy people encounters 4-5 times. Moreover, detecting hydrogen peroxide in EBC of pulmonary patients it is possible to monitor their therapy. As seen in fig. 1, exacerbation of asthma and its remission are characterized by statistically different levels of hydrogen peroxide concentration: remission causes the decrease of the latter.

We conclude that inflammatory pathologies cause the significant increase of hydrogen peroxide concentration in EBC pointing to pulmonary oxidative stress. It’s hence of great importance to measure a content of reducing marker.

Concentration of nitrite ion (NO_{2}^{-}) in EBC was found to be at the level of 10 µmol L^{-1} for healthy human (fig. 2). Micro-Molar content of nitrite is in accordance with the determined concentration of hydrogen peroxide in EBC. For low-inflammatory decease like COPD nitrite appears in EBC in quite similar concentrations. Exhaled breath condensate of the patients suffering from asthma contains statistically valuable reduced concentrations of NO_{2}^{-}.

Hence, nitrite content in EBC has an inversed correlation with hydrogen peroxide content: there is no significant difference between healthy human and low-inflammatory patients, where as in cases of inflammatory pathologies nitrite concentration in EBC is significantly decreased.
Figure 2. Concentration of nitrite ($\text{NO}_2^-$) in EBC of patients with difference pulmonary deceases

4. Conclusion

We conclude, that inflammatory pathologies cause a valuable increase of hydrogen peroxide coupled with a valuable decrease of nitrite concentrations in EBC. A simultaneous increase of oxidant and a decrease of reductant concentrations point to a significant pulmonary oxidative stress accompanying inflammatory pathologies. Curing of such pathologies can be, hence, improved using antioxidants.

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