Identification of nitric oxide in mitochondria of myometrium cell

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Aim. To demonstrate the possibility of NO synthesis in intact myocytes of uterus. Methods. Confocal scanning microscopy method, NO-sensitive fluorescent probe DAF-FM, MitoTracker Orange CM-H₂TMRos. Results. The basal production of NO in intact myocytes was shown using DAF-FM. Incubation of myocytes with NO donor – sodium nitroprusside (SNP) – led to an increase of the DAF-FM fluorescent signal. On the contrary, the addition of NO-synthase inhibitor – N-nitro-L-arginine (NA) – results in the reduction of fluorescent intensity. It was demonstrated colocalization of specific probe for mitochondria MitoTracker Orange CM-H₂TMRos and NO-sensitive dye DAF-FM. Conclusions. For the first time it has been demonstrated the presence of NO in smooth muscle cell mitochondria using laser confocal microscopy, NO-sensitive probe DAF-FM and specific marker of the functionally active mitochondria MitoTracker Orange CM-H₂TMRos.

Keywords: nitric oxide, myometrium, DAF-FM, confocal microscopy.

Introduction

Nitric oxide (NO) plays an essential role in many physiological processes, particularly in vasodilation, neurotransmission, immune responses, etc. [1, 2]. There are convincing proofs of the functional role of NO as a tocolytic agent. The ability of nitrocompounds to modulate activity of the Ca²⁺-transport system in myometrium cells was established [3]. At the same time, the identification of sources of its biosynthesis in the uterus myocytes is still a problem, because of a short time of existence and a high reactivity of nitric oxide.

The presence of the Ca²⁺-dependent isoforms of NO-synthase in mitochondria (mtNOS) was proved by immunohistochemical methods for single tissues [4–6]. On the one hand, nitric oxide can regulate the activity of electron transport chain in mitochondria through a reverse decrease in the cytochrome c-oxidase activity, and control of mitochondria pH. On the other hand, an excess production of NO together with intensification of the superoxide anion formation in mitochondria is accompanied by the peroxynitrite generation, damage of the respiratory chain components, mitochondria depolarization and the apoptosis development [4–6]. So in general, the mitochondria and cells life and destruction depend on the level of NO production. This indicates the importance of identification of NO biosynthesis in the intact cells mitochondria.

Modern fluorescent probe DAF-FM allows the recording of NO production, even the registration of the low NO concentrations (2–5 nM) in intact cells; the laser confocal microscopy method allows the visualization of the NO formation and the confirmation of the link of NO biosynthesis by these organelles using specific probe for mitochondria [4, 7].

So, the purpose of this work was to demonstrate the possibility of NO synthesis in the intact myocytes of uterus by energized mitochondria using the sensitive fluorescent dye DAF–FM, the selective...
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Materials and Methods

Myocytes were isolated from the uteruses of white nonlinear nonpregnant rats, by the method of Mollard using collagenase and soybean trypsin inhibitor [8]. The animals were narcotized by inhalation of diethyl ether, after that they were decapitated. All regulations for the work with laboratory animals were maintained (International Convention, Strasbourg, 1986).

Spatial distribution of fluorescent dyes in cells was investigated by the confocal laser scanning microscope LSM 510 META («Carl Zeiss», Germany). Myocytes were immobilized on poly-L-lysine. The cell immobilization, removal of unattached myocytes and all experimental procedures were conducted in physiological Henks’s solution (mM): NaCl – 136.9; KCl – 5.36; KH₂PO₄ – 0.44; NaHCO₃ – 0.26; Na₂HPO₄ – 0.26; CaCl₂ – 0.03; MgCl₂ – 0.4; MgSO₄ – 0.4; glucose – 5.5; Hepes (pH, 7.4, 37 °C) – 10 [9]. For myocytes permeabilization digitonin was added to the solution to a final concentration of 0.1 %.

For the visualization of mitochondria and cell nucleus the fluorescent dyes MitoTracker Orange CM-H₂TMRos (200 nM) and Hoechst 33342 (50 nM) were used respectively [10]. The loading of immobilized myocytes with NO-sensitive fluorescent probe DAF-FM (C₁₂H₁₄F₂N₂O₅, 4-amino-5-methylamino-2′,7′-difuorescin, diaminofluorescein-FM) in concentration 10 μM was carried out for 15 min at 24 °C. The experiments with confocal microscopy were performed in Multi Track mode. The Multi Track function permits several tracks to be defined as one configuration for the scan procedure. Each track is a separate unit and can be configured independently of the other tracks with regard to channels. Fluorescence of Hoechst 33342 was excited using laser wavelength 405 nm, and registered with the filter BP 505–530 [http://www.lifetechnologies.com/order/catalog/product/D23844, 9]. The study of fluorescent dye distribution in a cell was performed in Time Series mode. For the quantitative analysis of the time-dependent fluorescence intensity of the cell, the ROI (Region Of Interest) mode was used.

The data are presented as mean ± SEM [10].

In the work the following reagents were used: Hepes, glucose, saccharose, digitonin, sodium succinate, bovine serum albumin, poly-L-lysine, collagenase type IA, ATP, Pluronic F-27, DAF-FM, EGTA, CaCl₂ (Sigma, USA); DiOC₆(3), Hoechst 33342, soybean trypsin inhibitor (Fluka, Switzerland); MitoTracker Orange CM-H₂TMRos (Invitrogen, USA).

Results and Discussion

In the experiment an active acid form of DAF (DAF-FM) was used. It interacts directly with NO in the presence of O₂, forming triasole-fluorescein (DAF-FM-T) derivative, that has high quantum yield of fluorescence [7, 11]. The efficiency of DAF-FM entering into myoplasm was increased by permeabilization of the plasma membrane by 0.1 % digitonin, although the dye can penetrate partially through the membrane into a cell by diffusion [12]. The treatment of cells by the detergent significantly reduces the contribution of NO-synthase associated with the plasma membrane in a fluorescent signal. On the other hand, the dye entering the mitochondria was increased. These studies revealed (Fig. 1), that there was green fluorescent signal after pre-incubation with DAF-FM. The result reflects a basal NO production [4]. The signal was not associated with the cells autofluorescence in this area. A high sensitivity of the used dye permits to register a basal level of NO in myocytes, that is formed as a result of the constitutive NO-synthase functioning.

The signal is detected in the cell myoplasm, mapping its contours and forming the heterogeneous dyed areas. There were the NO-positively dyed areas also in the nuclei, and sometimes outside the cells, as a result of the diffusion.

The incubation of myocytes with 0.1 mM sodium nitroprusside (SNP) – NO donor – led to a substantial
Fig. 1. Distribution of DAF-FM-T (green color) and Hoechst 33342 (blue color) fluorescent signals in myocyte

Fig. 2. Increase in the intensity of fluorescent DAF-FM-T signal in myocytes after SNP addition; DAF-FM-T – green color, nucleus probe Hoechst 33342 – blue color

Fig. 3. Changes in fluorescent intensity of DAF-FM-T in myocytes with adding NA and SNP: MitoTracker Orange CM-H$_2$TMRos, red curve, channel 1; DAF-FM-T, green curve, channel 2; Hoechst 33342, blue curve, channel 3

Fig. 4. Quantitative analysis of DAF-FM-T fluorescent intensity changes in myocytes after adding NA and SNP relative to controls without these substances (100 %); $M \pm m, n = 3–7$

Fig. 5. Distribution of fluorescent dyes MitoTracker Orange CM-H$_2$TMRos (left figure, red color) and DAF-FM-T (right figure, green color) in myocyte

increase of fluorescent signal (Fig. 2). The fluorescence increased in myocytes and outside the cells, because NO was generated by SNP outside the cells too.

The incubation of cells with increased concentration (0.1 and 0.2 mM) of N-nitro-L-arginine (NA) – a nonselective inhibitor of the Ca$^{2+}$-dependent isoforms of NO-synthase [13] – was accompanied by the dose-dependent reduction of DAF-FM-T fluorescent intensity on average by 20 % (0.1 mM inhibitor) and 40 % (0.2 mM inhibitor). The growth of fluorescent response with adding SNP occurred in the presence of inhibitor (Fig. 3, 4).

These results show that DAF-FM responds to NO. DAF-FM does not test a level of a wide range of the active nitrogen and oxygen metabolites, for example NO$_2^-$, NO$_3^-$, ONOO$^-$, •O$_2^-$, H$_2$O$_2$, etc. [4]. There are good reasons to use DAF-FM for detection of nitric oxide in intact cell in our further investigations.

Colocalization of the NO-sensitive dye DAF-FM
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and the specific probe for mitochondria MitoTracker Orange CM-H$_2$TMRos, accumulated only in the intact energized organells [14], was demonstrated by the next studies (Fig. 5).

The computer analysis the image outside the cell nucleus has revealed an identical distribution of both fluorescent dyes (Fig. 6).

We have shown the potential presence of NO in mitochondria of uterus myocytes. Nonetheless, the nitric oxide biosynthesis in cells is linked not only with mitochondria. Constitutive NO-synthase is associated with the plasma membrane, sarcoplasmic reticulum and other intracellular structures. NO, synthesized by these structures, can diffuse to mitochondria and vice versa. Further researches are necessary to clarify the subcellular distribution of NO-synthases in the intact cells.

**Conclusions**

In this work for the first time the presence of NO in smooth muscle cell mitochondria has been demonstrated using the laser confocal microscopy, NO-sensitive probe DAF-FM and specific marker for the functionally active mitochondria MitoTracker Orange CM-H$_2$TMRos. It has been also shown that the amount of NO changes in the presence of the nitric oxide donor and NO-synthase inhibitor.

**REFERENCES**

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 можливість синтезу NO в інтактних міоцитах матки крыс. 
Методи. Конфокальна сканувальна мікроскопія, NO-чувствительний флуоресцентний зонд DAF-FM, MitoTracker Orange CM-H,TMRos. Результати. Показана базальна продукція NO в інтактних міоцитах за допомогою флуоресцентного зонду DAF-FM. Внаслідок інкубації клітин з донором NO нітропруссидом натрію (SNP) спостерігалось зростання флуоресцентного сигналу DAF-FM-T. І нарешті, внесення інгібітора NO-синтази – N-нітро-L-аргінину (NA) – призводило до гасіння флуоресценції. Доведено колокалізацію флуоресцентного мітохондріального

маркера MitoTracker Orange CM-H,TMRos та NO-чутливого зонду DAF-FM. Висновки. Вперше показано наявність NO в гладком’язових клітинах матки, і зокрема в мітохондріях, залученням методу лазерної конфокальної мікроскопії, NO-чутливого зонду DAF-FM та специфічного маркера функціонально активних мітохондрій MitoTracker Orange CM-H,TMRos.

Ключові слова: оксид азоту, міометрій, DAF-FM, конфокальна мікроскопія.

Идентификация оксида азота в митохондриях миоцитов матки
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Цель. Продемонстрировать возможность синтеза NO в интактных миоцитах матки крыс. Методы. Конфокальная сканирующая микроскопия, NO-чувствительный флуоресцентный зонд DAF-FM, MitoTracker Orange CM-H,TMRos. Результаты. Показана базальная продукция NO в интактных миоцитах с использованием флуоресцентного зонда DAF-FM. Инкубация миоцитов с донором NO — нитропруссидом натрия (SNP) — приводила к повышению флуоресцентного сигнала DAF-FM-T. И наоборот, внесение ингибитора NO-синтазы — N-нитро-L-аргинина (NA) — снижало интенсивность флуоресценции. Показана колокализация флуоресцентного митохондриального маркера MitoTracker Orange CM-H,TMRos и NO-чувствительного зонда DAF-FM. Выводы. Впервые показано присутствие NO в гладкомышечных клетках матки, в частности митохондриях, с использованием метода лазерной конфокальной микроскопии, NO-чувствительного зонда DAF-FM и специфического маркера функционально активных митохондрий MitoTracker Orange CM-H,TMRos.

Ключевые слова: оксид азота, миometрий, DAF-FM, конфокальная микроскопия.

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