Exposure to hyperbaric oxygen (HBO) can lead to seizures. Many studies have demonstrated that there exist a very close relationship between the alteration of cerebral blood flow (CBF) and the onset of seizures. Nitric oxide (NO) may play a key role in the change of CBF during exposure, and modulation of endothelial nitric oxide synthase (eNOS)-derived NO by HBO is responsible for early vasoconstriction, whereas late HBO-induced vasodilation depends upon a large amount of NO from both eNOS and neuronal nitric oxide synthase (nNOS). To investigate the effect of HBO on the activity and expression of eNOS in cerebral microvascular endothelial cells (CMEC) in vitro, primarily cultured CMEC from neonatal rats were exposed to oxygen at 500 kPa [5 atmosphere absolute (ATA)] for 10, 20, 30, 60 and 120 minutes (min), then eNOS activity, protein and mRNA contents in cells were detected. Our results showed that immediately after exposure, 30, 60 and 120 min HBO exposures did not alter NOS activity. When detected no matter immediately or six hours (h) after exposure, these exposures also did not alter eNOS protein and mRNA levels. However, when detected 24 h after exposure, 30, 60 and 120 min exposures upregulated eNOS protein content by 39%, 60% and 40% respectively. 10 and 20 min exposures upregulated eNOS mRNA content by about 15%, while 30, 60 and 120 min exposures upregulated it by about 20–30%. The increased eNOS protein and mRNA contents at 24 h after exposure may reflect new protein synthesis for eNOS. Our studies showed that with the exposing protocols we used, HBO did induce eNOS expression increase in CMEC. However, compared with the decrease of CBF in vivo, which occurred in a relative short time after rat was exposed to HBO above 4 ATA, the responses of eNOS in CMEC in vitro were a little slow. Thus we considered that for the vasodilation in the late period of HBO exposure before seizure, the effect of NO produced by eNOS was limited.

Introduction

Prolonged exposure to hyperbaric oxygen (HBO) is essential in some clinical and operational situations and can result in central nervous system (CNS) and pulmonary oxygen toxicity. The neurological manifestations of HBO toxicity include a variety of signs and symptoms ranging from anxiety, nausea, altered vision and hearing to muscular twitching, unconsciousness and convulsions. It has been reported that during HBO exposure, cerebral blood flow (CBF) decreased initially, then increased well before seizure onset.1-4 Therefore CBF increase could act as an objective pre-convulsive index and the overproduction of nitric oxide played an important role in such increase. The studies by Atochin et al. using endothelial nitric oxide synthase (eNOS)-/- and neuronal nitric oxide synthase (nNOS)-/- mutant mice suggested that eNOS derived nitric oxide (NO) was responsible for the early vasoconstriction responses, whereas late HBO-induced vasodilation depended upon both eNOS and nNOS.5 On the other hand, Hagioka et al. used selective nNOS inhibitor (7-NI) and the nonselective NOS inhibitor (L-NAME) elucidated the origins of NO production. Their data suggested that nNOS was the main mediator of NO production associated with increase in CBF during HBO exposure.6 To examine the exact inducible factor of HBO-induced vasodilation will be helpful to elucidate the mechanism of oxygen convulsion and to provide novel thoughts against this illness.

In this study, we investigated the direct effects of HBO exposure on primarily cultured rat cerebral microvascular endothelial cells (CMEC). After CMEC were exposed to 5 atmosphere absolute (ATA) 100% O2 for diverse exposure and sequentially cultured periods, eNOS activity, protein and mRNA levels in CMEC were detected.
Results

Effect of HBO exposure on the morphology of cultured endothelial cells. With an HBO exposure period of 120 minutes (min), the morphology of CMEC had no notable changes, no matter observed immediately after exposure or after the cells was sequentially cultured in 95% air—5% CO2 surroundings for 24 hours (h). The cells kept fusiform shape as that before exposure. Neighboring cells tightly attached to each other. Cells adhered to the culture dishes tightly. No notable decrease of cell quantity was observed (Fig. 1).

Effect of HBO exposure on viability of cultured endothelial cells. When detected 24 h after exposure, we found the viabilities of CMEC exposed to HBO and hyperbaric air (HBA) for 60 min were significantly higher than that of CMEC exposed to HBO for 120 min and normobaric air (NBA) (p < 0.01). There was no significant difference between the viabilities of CMEC exposed to HBO and that of CMEC exposed to HBA for 60 min. Also no significant difference existed between the viabilities of CMEC exposed to HBO for 120 min and that of CMEC exposed to NBA (p > 0.05) (Fig. 2).

Effect of HBO exposure on NOS activity in cultured endothelial cells. Nitric oxide synthase (NOS) activity in CMEC was measured immediately after exposure. Compared with NOS activity in CMEC exposed to NBA, activity in CMEC exposed to HBA for 60 min was not changed significantly. Similarly, activities in CMEC exposed to HBO for 30, 60 and 120 min were kept unchanged (Fig. 3).

Effect of HBO exposure on eNOS protein level in cultured endothelial cells. Light microscopic immunocytochemistry showed that eNOS was expressed in confluent endothelial cells in culture. eNOS was localized mainly in a paranuclear position, especially on one side of the nucleus (Fig. 4A and B). When compared no matter immediately or 6 h after exposure, there were no significant differences among the color intensities of stained eNOS protein in CMEC exposed to NBA, HBA and HBO for 60 and 120 min (Fig. 4A and B).

Western blotting analysis also indicated that eNOS protein contents in CMEC exposed to HBO for 60 and 120 min were not different from those in CMEC exposed to NBA or to HBA for 60 min, when detected no matter immediately or 6 h after exposure (p > 0.05) (Fig. 5A and B).

When measured 24 h after exposure, compared with eNOS protein contents in CMEC exposed to NBA, those in CMEC exposed to HBO for 10 and 20 min kept unchangeable (p > 0.05). However, as the exposure period prolonged, the eNOS protein contents increased. Figure 5C and D show 30, 60 and 120 min HBO exposure could increase eNOS contents by 39%, 60% and 40% respectively (p < 0.01) (Fig. 5C and D).

Effect of HBO exposure on eNOS mRNA level in cultured endothelial cells. When detected 24 h after exposure, RT-PCR analysis showed that eNOS mRNA contents in CMEC exposed to HBO for 60 and 120 min were not different from those in CMEC exposed to NBA or to HBA for 60 min (p > 0.05) (Fig. 6A and B).

However, when measured 24 h after exposure, compared with eNOS mRNA contents in CMEC exposed to NBA, 10 and 20 min
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enough for cell survival. Meanwhile, oxygen partial pressure in medium could reach to the predetermined level within a quite short time (5 ATA within 20 min), and this would be very favorable for investigations of HBO effects on cultured cells.

Based on the exposure methods, we found 120 min exposure to 5 ATA HBO produced no marked influences on the morphology and the quantity of CMEC when observed immediately after exposure. Even when observed at 24 h after exposure, the status remained unchangeable. The results of cell viability showed that there were no notable differences among the influences on cell viability of diverse exposure, although 60 min exposure could slightly increase cell viability. Meanwhile, we found such slight increase mainly resulted from the effect of high pressure, but not merely from HBO, as there was no significant difference between the result of 60 min HBO and 60 min HBA exposure. Summarily, we considered the exposure protocols we used had no notable influences on the living status of CMEC in certain period. On the base of such results, within certain period, the effects of HBO on the activity or content of a given enzyme or protein in CMEC would be direct and specific, and the potential changes of activity or content by HBO exposure would not be interfered by the living status of cells.

eNOS expression has been demonstrated in various types of endothelial cells, including CMEC. NO produced by CMEC may play a key role in the regulation of focal CBF under basal conditions.13 Studies in vivo had shown hyperoxia or hyperbaric oxygen exposure could change NO level in rat brain through modulating the activity and expression of eNOS and nNOS. But it was different on the point that which isoform of NOS is the main mediator. Thom et al. reported 2.8 ATA HBO exposure could increase NO concentration in rat cerebral cortex.4 Cerebral cortex blood flow increased during exposure and may be causally related to elevations of steady state NO concentration. Data showed nNOS activity contributes over 90% to total NO elevation. One important mechanism for nNOS activation was that HBO could trigger an increased association between nNOS and heat-shock protein (Hsp) 90, and subsequently increases intracellular Ca2+. Elayan et al. also reported a six- and fourfold increase in NO levels in the hippocampus and the striatum, respectively, of rats treated with 100% O2 at 3 ATA for two hours.14 Through determining neuronal NOS O2 Km, and using this value to estimate the effect of increasing brain O2 levels on NO production by nNOS, they found NO production by NOS closely matching the measured increase in vivo, indicating nNOS was mainly responsible for cerebral NO elevation under HBO conditions.

Studies on the relations among NO, oxygen free radicals and CBF revealed that HBO could cause a decrease in CBF in short period exposure. Such decrease was resulted from enhancing endogenous •O2¯ generation by hyperoxia, and because •O2¯ was a rapid neutralizer of NO, NO and its basal vasodilator effects were decreased.
As a proof, superoxide dismutase (SOD) could suppress •O$_2^-$ and increase CBF. Further investigations using genetically altered rodents showed NO neutralized by •O$_2^-$ and responsible for cerebral vascular constriction during breathing hyperbaric oxygen mainly consisted of eNOS-dependent NO. It was reasonable as the vascular endothelial cell and eNOS in it were the primary targets acted on by hyperoxia when high pressure oxygen was transported to brain.

As the exposure prolonged, although large numbers of oxygen free radicals were generated unceasingly, production of NO seemed to be more rapid due to the participation of abundant nNOS activated by HBO, exhibiting as a marked rise of CBF in the later period of a continuous HBO exposure, which always predicated a subsequent O$_2$ seizures. However, it was different on the point that which NOS isoform played a more important role in the modulation of the later vasodilation. Some considered NO by eNOS was responsible for the development of hyperoxic hyperemia preceding O$_2$ seizures, whereas neuronal NO may mediate toxic effects of HBO mainly by its reaction with superoxide to generate the stronger oxidant, peroxynitrite. Others suggested late HBO-induced vasodilation depends upon NO produced by both eNOS and nNOS. Besides, it was concurrent that nNOS was the main mediator of NO production responsible for CBF increase.

In view of the multiple effect of HBO on cerebral blood vessel constriction and dilation, and seeing that the key role of NO produced by CMEC on cerebral blood vessel tension, we investigated the direct influence of HBO on CMEC, especially on the activity and expression of eNOS in it, by directly exposing CMEC to HBO. Although some studies reported HBO could increase the expression of eNOS in cultured endothelial cells, effect of HBO on CMEC and eNOS in it had not been reported before.

Our studies found the effects of 5 ATA HBO on eNOS in primarily cultured rat CMEC were as follow:

1. When detected shortly after exposure, 30, 60 and 120 min exposure had no significant influence on NOS activity;
2. When detected shortly and six h after exposure, 60 and 120 min exposure had no significant influences on eNOS protein and mRNA levels;
3. At 24 h after exposure, 30, 60 and 120 min exposures could increase the eNOS protein content by 39%, 60% and 40% respectively, mRNA content by 20–30%. Although 10 and 20 min exposures had no effect on protein content, they could increase mRNA content by 15%. The increased eNOS protein and mRNA contents at 24 h after exposure may reflect new protein synthesis for eNOS.

Based on our results, we thought HBO exposure could really increase the production of eNOS in cultured endothelial cells. At the same time, we thought that compared with the decrease of CBF in vivo, which occurred in a relative short time after rat was exposed to HBO above 4 ATA, the responses of eNOS in CMEC in vitro were a little slow, even in consideration of the time needed for the balance of oxygen pressure in the culture medium and in ambient gas (5 ATA within 20 min). Therefore, the effect of NO came from eNOS was limited for the vasodilation in the late period of HBO exposure before seizure. Some other studies had shown that the impacts of HBO exposure were potent on nNOS in rat brain, and nNOS was the main mediator of NO production associated with increase in CBF during HBO exposure. Moreover, eNOS may not be inhibited in the early period of exposure, although CBF decreased rapidly after the oxygen pressure was raised. As suggested by Demchenko et al. and Zhilyaev et al. this transient decrease was mainly resulted from inactivation of eNOS-dependent NO by •O$_2^-$, decreasing the basal vasorelaxing action of NO.

Cerebral microvascular endothelial cell and neuron are different type cells. Our further investigations will focus on the difference of HBO modulation on NO production by eNOS in cerebral microvascular endothelial cell and by nNOS in neuron respectively.

Materials and Methods

Primary culture of rat CMEC. All experimental procedures were performed in adherence to the guidelines of Second Military Medical University on Animal Care and were approved by Second Military Medical University Animal Protocol Management and Review Committee.

Primary cultures of rat cerebral microvascular endothelial cells were prepared by modified methods described by Kis et al. and Abbott et al. Briefly, ten two-week-old SD rats were used for per culture. The cortices were minced and incubated in collagenase type II. Then bovine serum albumin/dulbecco's minimum essential
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medium (DMEM) was added to the homogenate and centrifuged. The pellet was washed and further digested in collagenase/dispase. The cell suspension was layered on a 50% Percoll gradient, and then centrifuged. The band of the endothelial cell clusters was aspirated and washed. The cells were seeded onto collagen type IV and fibronectin coated plastic dishes (for Western blotting and RT-PCR) or glass coverslip for immunohistochemistry. From the second day basic fibroblast growth factor was added. The culture medium was changed every other day. The cells reached a confluent monolayer on 5–7th day and consisted of more than 90% endothelial cells, verified by a spindle-shaped morphology and positive Factor VIII-related antigen. Such cells, after being starved with DMEM medium containing 1% fetal bovine serum (FBS) for 24 h, were used for all experiments.
HBO exposure. HBO exposure was performed in a small transparent chamber in which the temperature was controlled at 37 ± 0.5°C and the humidity was controlled at the level of 100 percent. The gas mixture was composed of 99% O₂ and 1% CO₂ in order to maintain physiological pH at the experimental pressure. Cells were treated in opened 35 mm culture dishes or plates containing appropriate volume of DMEM medium with 1% FBS. After flushed with mixed gas for two minutes, the chamber pressure was reached to 500 kPa (5 ATA) at rate of 100 kPa per min and maintained for 10, 20, 30, 60 or 120 min according to respective experiments. After exposure, cells were treated immediately or sequentially cultured in regular normobaric air incubator (NBA, 95% air—5% CO₂) for 6 or 24 h for various assays. Cells of negative and normal control were respectively exposed to HBA (4% O₂—95% N₂—1% CO₂) at 500 kPa and NBA for equivalent period as HBO exposure.

Morphological observation and viability assay. Cell morphology was assessed by means of a phase-contrast optical microscope (Olympus) immediately and 24 h after HBO exposure. Before viability assay, cells seeded on 96-well plates were exposed to HBA for 60 min, or exposed to HBO for 60 or 120 min. After exposure, cells were sequentially cultured in regular surroundings (95% air—5% CO₂) for 6 or 24 h. The viability was determined by the methyl thiazolyltetrazolium (MTT) assay. Briefly, MTT was added to the medium (final concentration 50 μg/ml) and cells were incubated for 4 h. After the crystals were completely dissolved in acidic isopropanol (0.01 M HCl in 5% isopropanol and 10% sodium dodecyl sulphate (SDS)) at 37°C, the optical density was determined at wavelength of 550 nm in ELISA reader (Bio-Tek Instruments). Results are expressed as mean ± standard deviation (SD). Statistical significance was determined by the Student’s t-test.

NOS catalytic activity. NOS activity was measured using Nitric Oxide Synthase Assay System (Beyotime, China), which is a simple and specific assay for the measurement of NOS activity in living cells under physiological conditions. This system use a cell-permeable fluorescent probe, 3-amino-4-aminomethyl-2',7'-difluorescein diacetate (DAF-FM DA), to react with NO produced by NOS to form a fluorescent triazolofluorescein. After exposure, cell monolayers were rinsed two times with 200 μl Hank’s Buffered Salt Solution (HBSS) each rinse to remove culture medium. Then 100 μl reaction buffer and 100 μl reaction mix containing the cofactor NADPH, arginine, and fluorescent probe were added to each well. Blank control was set by being only reaction buffer and reaction mix in well and cell-free. Incubate the plate at 37°C, protected from light for two hours. Read the plate in spectrofluorometer with an excitation filter at 495 nm and an emission filter at 515 nm. NOS activity was calculated by subtracting the value of blank control from the value of sample. The final representation of NOS activities in HBA and HBO exposed cells were presented as a percentage of that in NBA exposed cells.

Immunohistochemistry. Cells cultured on glass coverslip were fixed in 4% paraformaldehyde for 30 min immediately or six hours after HBO exposure. After pretreated in 1% hydrogen peroxide for 20 min, cells were incubated with rabbit polyclonal antibody against eNOS (1:100, Santa Cruz) for 24 h at 4°C. Then cells were incubated with Biotin-conjugated goat anti-rabbit antibodies (1:200) for 3–4 h at room temperature. Diaminobenzidin (DAB) was used as the chromogen.

Western blotting analysis. Cells were washed and incubated in lysis buffer, then harvested and briefly sonicated. After centrifugation, the supernatant containing the equal amount of protein was resuspended in loading buffer. The buffer was subjected to SDS-polyacrylamide gel electrophoresis and then transferred to polyvinylidenedifluoride (PVDF) membrane. The membrane was blocked with nonfat dry milk in buffer. Blocked membranes were incubated with primary antibody (polycolonal antibody to eNOS, and monoclonal antibody to β-actin, Santa Cruz) overnight at 4°C. Then membranes were incubated with 1:4,000 dilution secondary antibody (Santa Cruz) for one hour at room temperature. Proteins on the membranes were detected using an enhanced chemiluminescence detection kit (Amersham). For quantification, all blots were digitized with a scanner. Optical density of the bands was measured using Smartscape image analysis system (China). eNOS protein level was expressed as a ratio of eNOS’s bend densitometric value divided by that of β-actin’s respectively, and the final results were expressed as relative value referring to NBA eNOS protein level.

RT-PCR. Ice-cold lysis buffer was added to disrupt the cells. Total RNA isolation, reverse transcription, and semiquantitative competitive PCR were performed according to the instructions of RT-PCR kit (Qiagen). The sense (5’-TCC AGT AAC GAC AGT GCA-3’) and antisense (5’-CAG GAA GTA AGT GAG ACC-3’) primers for eNOS amplified a 693-bp fragment of murine eNOS that was confirmed by DNA sequencing. Each PCR cycle consisted of denaturing at 94°C for 30 seconds (s), annealing at 58°C for 30 s, and elongation at 72°C for 90 s. The linear exponential phases for eNOS and glyceraldehydes phosphate dehydrogease (GAPDH) PCR were all 30 cycles. Equal amounts of corresponding GAPDH reverse transcription-PCR products were loaded on 1.5% agarose gels. Optical densities of ethidium-bromide stained DNA bands were quantitated using Smartscape image analysis system (China). eNOS mRNA level was expressed as a ratio of eNOS’s bend densitometric value divided by that of GAPDH’s respectively, and the final results were expressed as relative value referring to NBA eNOS mRNA level.

Statistical analysis. Data are presented as mean ± SD. Differences between treatment and control groups were compared by unpaired two-tailed Student’s t-test or by ANOVA. p values of <0.05 were considered statistically significant.

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