Dissociation of DNA damage and mitochondrial injury caused by hydrogen peroxide in SV-40 transformed lung epithelial cells

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Abstract

Background: Since lung epithelial cells are constantly being exposed to reactive oxygen intermediates (ROIs), the alveolar surface is a major site of oxidative stress, and each cell type may respond differently to oxidative stress. We compared the extent of oxidative DNA damage with that of mitochondrial injury in lung epithelial cells at the single cell level.

Result: DNA damage and mitochondrial injury were measured after oxidative stress in the SV-40 transformed lung epithelial cell line challenged with hydrogen peroxide (H₂O₂). Single cell analysis of DNA damage was determined by assessing the number of 8-oxo-2-deoxyguanosine (8-oxo-dG) positive cells, a marker of DNA modification, and the length of a comet tail. Mitochondrial membrane potential, ∆Ψm, was determined using JC-1. A 1 h pulse of H₂O₂ induced small amounts of apoptosis (3%). 8-oxo-dG-positive cells and the length of the comet tail increased within 1 h of exposure to H₂O₂. The number of cells with reduced ∆Ψm increased after the addition of H₂O₂ in a concentration-dependent manner. In spite of a continual loss of ∆Ψm, DNA fragmentation was reduced 2 h after exposure to H₂O₂.

Conclusion: The data suggest that SV-40 transformed lung epithelial cells are resistant to oxidative stress, showing that DNA damage can be dissociated from mitochondrial injury.

Background

Aerobic cells are constantly exposed to reactive oxygen intermediates (ROIs). Increased intracellular levels of the ROIs, superoxide (O₂⁻), hydroxyl radical (·OH), or hydrogen peroxide (H₂O₂) are referred to as oxidative stress [1,2]. The alveolar surface of the lung is a major target for oxidative stress. Also, cigarette smoking can cause an acute inflammatory reaction in the lung, characterized by the accumulation and activation of leukocytes in vivo, producing ROIs and nitrogen species in high concentrations [3,4]. These different reactive species might be responsible for much of the tissue injuries and disease states associated with inflammation. Increased oxidative DNA damage, such as 8-hydroxyguanine formation in human lung tis-
sue [5], is found in smokers compared with non-smokers. Oxidative stress may play an important role in the pathogenesis of smoking-associated diseases, such as chronic obstructive pulmonary diseases (COPD) [6,7], asthma [8], and carcinogenesis [9–11].

ROIs damage DNA, resulting in base modifications, e.g. 8-oxo-2′-deoxyguanosine (8-oxo-dG) [12,13], as well as causing DNA strand-breaks [14]. The presence of 8-oxo-dG in DNA is considered as a marker of oxidative stress and DNA damage [13]. We quantified DNA modifications induced by H₂O₂, as shown by 8-oxo-dG-positive cells at the single cell level, using laser-scanning cytometry (LSC). In general, this method of analysis can be applied to stained cells adherent to slides, eliminating the cell loss that inevitably occurs due to repeated centrifugations during sample preparation for flow cytometry. And also, the alkaline version of the single cell gel electrophoresis (SCGE) or Comet assay represents a sensitive technique for the detection of single-stranded DNA breaks [15]. We used the Comet assay to measure the extent of single-stranded DNA breaks in epithelial cells treated with H₂O₂, and compared this with the number of base modifications at the single cell level.

An altered mitochondrial membrane potential followed by increased ROI generation, the loss of mitochondrial cardiolipin, and increased intracellular Ca²⁺ concentrations have recently been described as common features of apoptosis [16]. Apoptosis is a programmed event, which is regulated by genes, and is irreversible in terms of the DNA cleavage.

The aim of the current study was to investigate the susceptibility of DNA damage and mitochondrial injury caused by H₂O₂ in cultured SV-40 transformed lung epithelial cells, which may prove a useful in vitro model of the lung. The extent of oxidative DNA damage and mitochondrial injury was assessed at the single cell level. We hypothesized that lung epithelial cells would be relatively resistant to oxidative stress and one may therefore be able to distinguish between DNA damage and mitochondrial injury.

**Results**

**Cell cycle analysis**

To compare the cell cycle effects of different culture conditions on starved BEAS-2B cell lines treated with H₂O₂, the number of proliferating cells (defined as BrdU-positive cells) and apoptotic cells were measured after culturing with growth factors (GF) or in GF-free medium. Quiescent cells starved of GF for 48 h were exposed to H₂O₂ (100–500 μM) for 1 h, after which the cells were cultured in fresh media without GF. After 1 day, cycle analysis indicated that the cells treated with H₂O₂ were similar to untreated controls (Fig. 1A and 1C). However, when quiescent cells were exposed to H₂O₂ followed by culturing in fresh medium containing GF to initiate entry into cycle, a higher percentage of G₁ phase-cells (83%) and a lower percentage of S phase-cells (6%) were found compared to the controls (70% and 16% respectively; Fig. 1B and 1C). H₂O₂ inhibited cell proliferation in a concentration-dependent manner in conjunction with GF (Fig. 1C). As a result, few apoptotic cells (3%) were detected, seen as a "pre-G₁" phase population (Fig. 1B).

**8-oxo-dG for detecting DNA damage**

It had previously been shown by high-performance liquid chromatography (HPLC) that H₂O₂ induces 8-oxo-dG, a DNA damage marker [16]. Exposure of quiescent WI-26VA4 cells, another SV-40 transformed epithelial cell line, to H₂O₂ had no effect on the number of 8-oxo-dG-positive cells (Fig. 2A). In the present study, we used a direct binding fluorescent probe to detect 8-oxo-dG in the DNA, which was quantified by LSC. The rate of DNA damage measured was that induced by a 1 h exposure of BEAS-2B cells to H₂O₂. We also assessed the number of 8-oxo-dG-positive cells by LSC (Fig. 2B). H₂O₂ treatment increased the number of 8-oxo-dG-positive cells increased in a concentration-dependent manner (2.1 versus 19.6%; Fig. 2C).

**Comet assay**

To determine the effect of H₂O₂ on single-strand DNA breaks in BEAS-2B cells, cells were treated with H₂O₂ (200–1000 μM) and the tail moment (TM) determined. The mean tail moment relative to time of H₂O₂ treatment is shown in Figure 3B. For the first 1 h after H₂O₂ treatment, the mean TM showed both a time- and concentration-dependent decrease. However, the mean TM increased at every concentration of H₂O₂ between 60 and 120 min.

**JC-I staining**

Previous studies have suggested that a decline of ΔΨₘ may be an early event in the process of cell death, and so we determined ΔΨₘ at various times after H₂O₂ treatment in BEAS-2B cells, using the membrane potential-sensitive probe, JC-1, which forms monomers (green fluorescence) at a low membrane potential and J-aggregates (red fluorescence) at a higher membrane potential [17]. The ratio between the red and the green signals is indicative of the ΔΨₘ. There was a dramatic drop of the red fluorescence in cells exposed to H₂O₂, indicating a loss of ΔΨₘ. Figure 4A shows a clear increase in the percentage of cells that emitted only green fluorescence after H₂O₂ treatment, representing cells with a depolarized mitochondrial membrane. H₂O₂ caused a concentration-dependent reduction or loss in ΔΨₘ (Fig. 4B).
Figure 1
Distribution of cell cycle phases in BEAS-2B cells exposed to H₂O₂. A, quiescent cells were exposed to H₂O₂ for 1 h, followed by culturing in growth factor (GF) free-media for 24 h. B, quiescent cells were exposed to H₂O₂ for 1 h, followed by culturing in media supplemented with GF for 24 h. All floating and attached cells were collected, incubated with BrdU, and stained with anti-BrdU and PI. C, graphic representation of H₂O₂-exposed BrdU-positive cells. *, significantly different from control at p < 0.05 (Mann-Whitney U test).
Figure 2
A, green-fluorescent image of intranuclear 8-oxoguanine expression. WI-26VA4 cells were cultured with and without 1 mM H$_2$O$_2$ for 1 h. The cells were fixed with cold methanol in situ, then stained with a protein, which binds specifically and directly to 8-oxo-dG. B, representative dot plot of DNA content vs. 8-oxo-dG of FITC staining in WI-26VA4 cells treated with or without 1 mM H$_2$O$_2$ for 1 h. Cells were counterstained with PI, and analyzed by laser scanning cytometer. The number in each dot-plot indicates the percentage of 8-oxo-dG positive cells. C, graphic representation of the 8-oxo-dG positive cells that increased in an H$_2$O$_2$ concentration-dependent manner. *, **, significantly different from controls at $p < 0.05$ and $p < 0.01$, respectively (Mann-Whitney U-test).
Figure 3
A, typical DNA migration patterns of BEAS-2B cells treated with 500 µM H₂O₂ for 1 h at 37°C. B, tail moment (TM) representing the DNA breaks induced by H₂O₂ in BEAS-2B cells. Cells were exposed to H₂O₂ at 37°C for 1 h and then subjected to alkaline single-cell gel electrophoresis (Comet assay). Values are expressed as the means ± SD of 3 experiments carried out in duplicate (n = 3). *, **, significantly different from controls at p < 0.05 and p < 0.01, respectively (Mann-Whitney U test).
Figure 4
A. Flow cytometric analysis of the preconditioning-induced changes in the inner mitochondrial membrane potential in BEAS-2B cells. The BEAS-2B cells were exposed to H₂O₂, and stained with the potentiometric dyes JC-1. Bivariate plots of red (FL2) versus green (FL1) fluorescence were used as an estimate of mitochondrial membrane potential (ΔΨₙ). B, percent of cells with ΔΨₙ disruption vs. exposure time. Cells were treated with 200, 500 µM, and 1 mM of H₂O₂ for 10, 30, 60, and 120 min. Untreated controls and H₂O₂-treated cells were then stained with JC-1 and analyzed by flow cytometry as described in Methods. n = 10,000 cells/analysis. *, **, significantly different from controls at p < 0.05 and p < 0.01, respectively (Mann-Whitney U-test).
Immunohistochemistry for nitrotyrosine

An important pathway that enhances NO cytotoxicity is the rapid reaction with superoxide anion radicals (O₂⁻) to yield peroxynitrite (ONOO⁻), which has a wide range of biologic activities that includes oxidation of biomolecules and protein tyrosine nitration [18]. It is widely accepted that peroxynitrite is a key mediator of tissue injury in oxidative stress. To determine whether H₂O₂-induced peroxynitrite formation occurred in BEAS-2B cells, cells were stained with an anti-nitrotyrosine antibody. H₂O₂ (60 min, 500–1000 µM) induced clear perinuclear nitrotyrosine staining (Fig. 5B and 5C). The peroxynitrite donor, SIN-1, induced a predominantly nuclear membrane distribution of nitrotyrosine staining (Fig. 5D).

Discussion

The alveolar surface of the lung is a major site of oxidant injury. In lung epithelial cells, oxidative damage to DNA is caused not only by air pollutants, but also by neutrophil products [19]. Using high-density DNA microarrays, lung epithelial cells exposed to oxidative injury showed early induction of bcl-2 and mdm-2, genes which are known to be are involved in the regulation of apoptosis [20]. Both DNA and mitochondrial events combine to regulate cell cycle progression and apoptosis after treatment with H₂O₂. This study has assessed DNA damage and mitochondrial injury at the single cell level.

First, we have confirmed an effect of H₂O₂ on replicating DNA and apoptosis during the cell cycle. Oxidative stress is believed to induce the cyclin dependent kinase inhibitor, p21, thereby inhibiting DNA replication and hence allowing additional time to repair damaged DNA. Consistent with a previous report [21] demonstrating that H₂O₂ induced G₁ arrest in A549 cells, we have shown that a 1-h pulse of H₂O₂ induced cell arrest at G₀/G₁ phase and small amounts of apoptosis (only 3%) in SV-40 transformed lung epithelial cells in the presence of growth factors.

A variety of endogenous and environmental factors can cause DNA damage [22,23]. Estimates indicate that a human cell sustains ~10,000 lesions per day due to metabolism-generated free radicals [23]. There are two types of DNA damage induced by the genotoxic substance H₂O₂, DNA modifications and DNA breaks. Over 100 oxidative stress-induced DNA modifications have been characterized [24,25], of which the 8-oxo-dG lesion is the most common and extensively studied oxidative lesion [26]. To locate of oxidative damage in situ, a number of different investigators have developed antibodies against the oxidative DNA lesion, 8-oxo-dG [27–29]. We used an FITC-conjugated substrate to detect 8-oxo-dG and analyzed this by LSC at the single cell level. Traditional methods using HPLC [16] can assess the total amount of DNA damage per genomes whereas LSC indicates DNA damage at each individual cell. The detection of 8-oxo-dG by fluorescence in a flow cytometer [30] or by LSC might be useful tools for estimating DNA damage at the single cell level. Consistent with Dizdaroglu et al. [26], we have shown that 8-oxo-dG accounts for 5% of the total oxidized bases known to occur in DNA, and that even untreated cells showed 8-oxo-dG positivity (2.1%; Fig. 2B). In addition, we have demonstrated that H₂O₂ induced 8-oxo-dG-positive cells appear in a concentration-dependent manner.

The Comet assay is a sensitive method for the detection of DNA strand-breaks and alkali-labile (e.g. apurinic) sites in cells exposed to genotoxic carcinogens [15]. In this study, we did not use a modified comet assay with a purified DNA repair enzyme, formamidopyrimidine DNA-glycosylase (FPG), even though this method is more sensitive than the assay for detecting FPG-sensitive sites than our classic comet assay [31,32]. However, FPG is one of the multifunctional DNA repair enzymes participating in base excision repair [33]. We used the classic comet assay to detect repair phase of DNA breaks. To evaluate DNA fragmentation as a consequence of oxidative stress, cells treated with H₂O₂ showed an initial concentration-dependent increase in strand-breaks up to 1 h before a subsequent decrease in DNA fragmentation, which may reflect the mechanism involved in the repair of damaged DNA.

The increasing amounts of 8-oxo-dG and tail moments following H₂O₂ exposure suggest that 8-oxo-dG might give an estimate of the levels of DNA fragmentation. This finding is disagrees with previous reports [34,35], which demonstrated significant increases in fragmented DNA in cells treated with H₂O₂ but no concomitant increase in 8-oxo-dG. Glutathione (GSH) is a major non-protein antioxidant that protects cells from oxidative stress. The airway epithelium is a target for ROS, either inhaled or released during lung inflammation, and is resistant to oxidant injury due to its abundant level of intracellular GSH [36]. Another explanation is that lung epithelial cells may be susceptible to repair DNA damage by FPG as well as 8-oxo-dG DNA glycosylase (OGG-1) [33]. In airway epithelial cells where DNA repair occurs, we see parallel induction of both DNA strand-breaks and DNA modifications, albeit both at low levels.

In this study, the possibility exits that resistance of the SV-40 transformed cell line to DNA damage may not be the same as that of human primary lung epithelium, because SV-40 large T antigen simultaneously disables the retinoblastoma (pRB) and p53 tumor suppressor pathways [37]. More experiments on other types of lung epithelial cells transformed by different means need to be analysed to establish the mechanism of DNA repair.
Recent progress has shown the central role of mitochondria in death signaling [38]. In certain situations, death stimuli induce an increase in the permeability of the outer mitochondrial membrane that releases apoptosis inducers, such as cytochrome c [39] and apoptosis-inducing factor (AIF) [40]. In our study, the $\Delta \Psi_m$ was quantified by the flow cytometric analysis of JC-1-stained cells. This potential is the driving force for crucial cellular functions, ranging from ATP synthesis to the participation of mitochondria in Ca$^{2+}$ signaling. We have shown that H$_2$O$_2$ causes a concentration-dependent dissipation of $\Delta \Psi_m$ (Fig. 4). Under conditions where extensive growth-arrest and moderate apoptosis were observed, others have reported a general degradation of mitochondrial, but not nuclear polynucleotides during the early stage response to oxidative stress [41]. Mitochondria are not only the major site for the generation of reactive oxygen species, but one of the main targets of oxidative damage. One of the major products of DNA oxidation, 8-oxo-dG, accumulates in mitochondrial DNA (mtDNA) to three times the level in nuclear DNA [42]. Mitochondrial DNA is particularly susceptible to damage by ROIs because of its close proximity to the electron transport chain and its lack of protective histones. Prolonged dissipation of $\Delta \Psi_m$ might be attributed to this susceptibility of mitochondrial DNA damage.

The involvement of peroxynitrite (ONOO$^-$) in inflammatory diseases has been implicated by the detection of 3-nitrotyrosine, an allegedly characteristic protein oxidation products.
product, in various inflamed tissues. The formation of reactive nitrogen species (RNS), including peroxynitrite and nitrogen dioxide (NO₂), may be the prime reason in many cases why NO· can contribute to the etiology of inflammatory lung diseases, such as asthma [43]. Detection of 3-nitrotyrosine at inflammatory sites serves as a biochemical marker for peroxynitrite formation [18,44]. Our results indicate that H₂O₂ and the SIN-1 might produce peroxynitrite, which was detected as 3-nitrotyrosine immunoreactivity. We suggest that H₂O₂, as well as peroxynitrite, might induce DNA damage and mitochondrial injury.

In conclusion, DNA modifications, detected as 8-oxo-dG positive cells, and DNA fragmentation, assessed by the comet tail length, showed a similar pattern and were increased in lung epithelial cell lines exposed to H₂O₂. H₂O₂ induced the cells to undergo apoptosis and/or G₁ arrest, which is the cell-cycle phase for repairing DNA damage. The nucleus and mitochondria of the SV-40 transformed lung epithelial cells exhibited different susceptibilities to oxidative stress injury. This may be the mechanism of apoptosis, which is a programmed event, and is irreversible in terms of DNA cleavage. The mechanism of lung resistance to oxidative stress may be in the dissociation of mitochondria damage from the cascade that induces apoptosis, thus increasing the threshold for DNA repair.

**Methods**

**Cell culture**

BEAS-2B cells, which are SV-40 immortalized cells derived from normal human bronchial epithelium [45], were obtained from American Type Culture Collection (Manassas, VA). The cells (passages 32 to 70) were cultured in serum-free keratinocyte basal medium (Sigma; St. Louis, MO), which was supplemented with 10 ng/ml of epidermal growth factor (Gibco BRL; Grand Island, NY) and 30 μg/ml of bovine pituitary extract (Gibco BRL) in the presence of 5% CO₂ at 37°C. The human lung cell line WI-26VA4 was kindly provided by Human Science Research Resource Bank (Osaka, Japan). Cells (passages 162 to 180) were cultured in Dulbecco’s modified Eagle’s medium (DMEM; IWAKI, Chiba, Japan) supplemented with 10% fetal calf serum, amino acids (L-arginine HCl 252.8 mg/l, L-cystine 48.0 mg/l, L-histidine HCl 83.8 mg/l, L-isoleucine 105.0 mg/l, L-leucine 105.0 mg/l, L-lysine HCl 146.2 mg/l, L-methionine 29.8 mg/l, L-phenylalanine 66.0 mg/l, L-threonine 95.2 mg/l, L-tryptophan 20.4 mg/l, L-tyrosine 72.4 mg/l, and L-valine 93.8 mg/l), mixed vitamins (folic acid 1.0 mg/ml, nicotinamide 1.0 mg/ml, pyridoxal HCl 1.0 mg/ml, riboflavin 0.1 mg/ml, and thiamin HCl 1.0 mg/ml), penicillin (100 U/ml), and streptomycin (100 μg/ml) in the presence of 5% CO₂ at 37°C.

**Cell cycle analysis using BrdU incorporation**

BEAS-2B cells exposed to H₂O₂ were pulse-labeled with 50 μM 5-bromo-2′-deoxyuridine (BrdU) (Sigma Aldrich; Tokyo, Japan) for 30 min one day after exposure, and harvested by trypsinization, pelleted, and gently resuspended in phosphate buffered saline (PBS; OXOID, Hampshire, England). The cells were fixed with ice-cold 70% ethanol, centrifuged and resuspended in 1 ml 2 N HCl containing 0.05% Triton X-100 (v/v). The cells were incubated at room temperature for 30 min followed by centrifugation, rinsed in 0.1 M borate buffer, pH 8.5, and PBS/bovine serum albumin (BSA; Sigma-Aldrich) (1 mg/ml BSA in PBS), after washing with PBS/0.5% Tween 20/1% BSA. Appropriately diluted anti-BrdU antibody (clone BU-1; Amersham, Buckinghamshire, UK) was added to the cell pellet. After incubation for 1 h at room temperature, the cells were rinsed twice in PBS/BSA. For visualization, fluorescein isothiocyanate (FITC)-conjugated Fab2 fragments of rabbit anti-mouse Ig (DAKO, Glostrup, Denmark) antibody were added in a 1:30 dilution. After incubation for 45 min at room temperature, the samples were rinsed twice in PBS/BSA and the cells were resuspended in 0.5 ml cold PBS supplemented with 100 μg/ml RNase (Sigma-Aldrich) and 20 μg/ml propidium iodide (PI; Sigma-Aldrich). The samples were allowed to stand for 15 min on ice in the dark before flow cytometric analysis. For the negative control, the primary antibody was omitted. The stained cells were analyzed by FACScan (Becton-Dickinson, Franklin Lakes, NJ), using the CELL QUEST program.

**Measurement of 8-oxo-2′-deoxyguanosine (8-oxo-dG) levels in DNA by laser scanning cytometer**

The extent of cell DNA damage was determined using a commercial kit (Calbiochem-Novabiochem Corporation; Darmstadt, Germany) on WI-26VA4 cells at 50 to 70% confluence on coverslips in a 6-well culture plate. The cells were fixed with fresh 4% paraformaldehyde (Wako Life Science, Osaka, Japan) in PBS and dehydrated in 70% ethanol, after washing with PBS/Tween 20, containing thimerosal. To block non-specific binding sites, a blocking solution was added to each well and incubated for 1 h at 37°C. The blocking solution was removed and the cells washed with PBS/Tween 20 before being incubated with FITC-conjugated substrate for 1 h at 37°C, and washed 5 times with PBS/Tween 20 with 3 min of gentle shaking per wash. Next, the cells were incubated with PI-RNase for 10 min at 37°C, washed 5 times with PBS/Tween 20 with 3 min of gentle shaking per wash, and rinsed once with distilled water. The coverslips were freed, mounting solution was added, and placed on slides to be observed under fluorescent microscopy with FITC filters.
Cells were subsequently analyzed with an LSC (LSC101; Olympus, Tokyo, Japan) as previously described [46], using a krypton/argon laser operating at 488 nm and 568 nm. Dual-channel imaging was performed with two photomultipliers, using a band-pass filter specifically designed to detect FITC fluorescence and a long-pass (590 nm) filter for detecting PI fluorescence. Images were generated using integral computational and display software. To assess mean fluorescence intensity (MFI) of 8-oxo-dG, each nucleus was identified by areas counterstained with PI. We analyzed a 10-mm² scan area, occupied by approximately ~1 × 10⁴ cells.

**Comet assay**

The alkaline version of the Comet assay was performed according to the method of Singh et al. (1988) with minor modifications [15]. Briefly, fully-frosted microscope slides were pre-coated with 1 ml of 0.75% normal melting point agarose (NMA; Promega, Madison, WI) and stored at 4°C. This layer was removed before use and 120 µl of 0.75% NMA was pipetted onto the slides, which were then covered with coverslips. Cell suspensions of BEAS-2B cells (1 × 10⁴ cells/10 µl) were mixed with 70 µl of 1% low melting point agarose (LMA) and pipetted over the first layer of agarose. NMA (80 µl) was used as a final protective layer. After each step, the slides were incubated at 4°C for 10 min to allow the agarose to gel. Slides were placed in cold lysing solution (2.5 M NaCl, 100 mM Na₂EDTA, 10 mM Tris, pH 10, and 1% sodium N-lauroyl sarcosinate to which 10% dimethyl sulfoxide and 1% Triton X-100 had been added immediately prior to use) for 1 h. After lysis, the slides were placed in electrophoresis buffer (300 mM NaOH and 1 mM Na₂EDTA, pH 13) for 20 min to allow unwinding of the DNA. Electrophoresis was conducted in the same buffer by applying an electric current of 0.8 V/cm (300 mA) for 20 min, using an electrophoresis power supply (Pharmacia LKB, Sweden). Finally, the slides were washed in neutralization buffer (0.4 M Tris, pH 7.5) 3 times for 5 min each, dried and stained with Hoechst 33342 (10 µg/ml).

Image analysis involved a microscope equipped with an LSC. Nuclei were excited with green light and the emitted red spectrum was captured by a 40× dry objective. Images of 50 randomly selected nuclei were analyzed per duplicate slide (100 nuclei in total). The tail moment (TM) was determined to assess the extent of DNA damage. The TM value was defined as the product of the percentage of DNA in the comet tail and the tail length.

**Determination of mitochondrial membrane potential (ΔΨm)**

BEAS-2B cells were stained with 1 µg/ml of 5,5',6,6'-tetrachloro-1,1',3,3'-tetracyano-9,9'-bianthrylcarbocyanine (JC-1; Molecular Probes, Eugene, OR) for 30 min at 37°C, trypsinized, resuspended in medium at a density of approximately 1 × 10⁷/ml, and transferred on ice to the flow cytometer. JC-1 was excited at 488 nm and the monomer signal (green) was analyzed at 525 nm (FL1) on a FACSscan (Becton-Dickinson, Mountain-View, CA) flow cytometer. Simultaneously, the aggregate signal (red) was analyzed at 590 nm (FL2). In BEAS-2B cells, the percentage of PI-negative cells (1 × 10⁴) was analyzed and the mean fluorescence was calculated for each sample.

**Immunohistochemistry for nitrotyrosine**

BEAS-2B cells were cultured in 6-well plates on coverslips and were untreated or treated with H₂O₂ alone or with a peroxynitrite generator 3-morpholinosyndonimine (SIN-1), which is the precursor of peroxynitrite, for 1 h. The coverslips were then washed with PBS, prefixed in 2% formalin at room temperature for 10 min, and fixed in methanol/acetone (1:1) at -20°C for 10 min. The coverslips were washed with PBS and then were blocked in PBS containing 20% normal swine serum (Dako), 0.4% Triton X and 0.1% BSA. The fixed cells were stained with a primary antibody against nitrotyrosine (Upstate Biotechnology, Lake Placid, NY) diluted at 1:50 in PBS containing 0.4% Triton X and 0.1% BSA for 1 h. After removing the first antibody by washing, the cells were incubated with biotinylated goat anti-rabbit Ig (Dako) in PBS containing 0.4% Triton X and 0.1% BSA for 45 min. After removing the second antibody, the cells were incubated with FITC-conjugated streptavidin (Dako) in PBS containing 0.4% Triton X and 0.1% BSA for 45 min. Finally, the cells were counter-stained with 20% hematoxylin and the coverslips were mounted on slides with mounting medium (DPX). The stained cells were visualized and photographed using a confocal laser scanning microscope (Leica, Heidelberg, Germany).

**Data analysis**

Group data are expressed as the mean (± SD). Comparisons between data were tested using the Mann-Whitney U test. A p value < 0.05 was considered significant.

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