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Asbestos Inhalation Induces Reactive Nitrogen Species and Nitrotyrosine Formation in the Lungs and Pleura of the Rat

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Abstract

To determine whether asbestos inhalation induces the formation of reactive nitrogen species, three groups of rats were exposed intermittently over 2 wk to either filtered room air (sham-exposed) or chrysotile or crocidolite asbestos fibers. The rats were killed at 1 or 6 wk after exposure. At 1 wk, significantly greater numbers of alveolar and pleural macrophages from asbestos-exposed rats than from sham-exposed rats demonstrated inducible nitric oxide synthase protein immunoreactivity. Alveolar macrophages from asbestos-exposed rats also generated significantly greater nitrite formation than did macrophages from sham-exposed rats. Strong immunoreactivity for nitrotyrosine, a marker of peroxynitrite formation, was evident in lungs from chrysotile- and crocidolite-exposed rats at 1 and 6 wk. Staining was most evident at alveolar duct bifurcations and within bronchiolar epithelium, alveolar macrophages, and the visceral and parietal pleural mesothelium. Lungs from sham-exposed rats demonstrated minimal immunoreactivity for nitrotyrosine. Significantly greater quantities of nitrotyrosine were detected by ELISA in lung extracts from asbestos-exposed rats than from sham-exposed rats. These findings suggest that asbestos inhalation can induce inducible nitric oxide synthase activation and peroxynitrite formation in vivo, and provide evidence of a possible alternative mechanism of asbestos-induced injury to that thought to be induced by Fenton reactions. (J. Clin. Invest. 1998. 102:445–454.) Key words: crocidolite • chrysotile • nitric oxide • inducible nitric oxide synthase • nitrotyrosine

Introduction

It is well-recognized that occupational asbestos exposure is causally linked to a variety of clinical disorders afflicting the lungs and serous membranes including asbestosis, parietal pleural plaques, visceral pleural fibrosis, bronchogenic carcinoma, and diffuse malignant mesothelioma of the pleura and peritoneum (1, 2). Although all commercial varieties of asbestos are capable of fibrogenic and carcinogenic insult, there has been a long-standing debate regarding the relative potency of chrysotile (the serpentine type of asbestos that has been used most extensively in North America) versus crocidolite or amosite (amphibole) asbestos in this regard (3–5). Considerable attention has been focused on the putative role of reactive oxygen species in mediating asbestos-induced injury to the lungs and pleura via superoxide anions (O₂⁻), hydrogen peroxide (H₂O₂), and hydroxyl radicals (-OH) generated by modified Haber-Weiss (Fenton) reactions. Several studies have implicated the generation of reactive oxygen species in crocidolite-induced cellular cytotoxicity (6–8) and demonstrated that crocidolite-induced pulmonary inflammation and fibrosis in the rat was abrogated by the administration of polyethylene-conjugated catalase via an osmotic pump (7).

While generation of -OH by Fenton reactions may explain some of the biologic effects of crocidolite (chemical composition: Na₂[Fe³⁺⁺][Fe²⁺⁺][SiO₄]²⁻[OH]₄), it cannot account for the cytotoxic actions of chrysotile (chemical composition: Mg₃[SiO₄]²⁻[OH]₄), which contains only 2–3% elemental iron (9, 10). Therefore, it is conceivable that asbestos fibers also may induce cellular injury via mechanisms not based on Fenton chemistry such as by the generation of reactive nitrogen species (RNS), as has been demonstrated in human acute lung injury and experimental models of granulomatous lung inflammation (11, 12). Some support for this contention is provided by studies which demonstrated that the addition of either crocidolite or chrysotile fibers to cytokine-stimulated cell cultures upregulated the production of nitric oxide (iNO) by rat alveolar macrophages (13) as well as by rat parietal pleural mesothelial cells (14). In the latter study, asbestos challenge also generated the formation of the highly reactive nitrogen compound, peroxynitrite anion (ONOO⁻), a finding that was associated with cytotoxic changes within the pleural mesothelial cells (14). Furthermore, crocidolite fibers have been shown to induce the formation of the potentially carcinogenic modified DNA base, 8-hydroxy-2'-deoxyguanosine, in A549 human lung epithelial cells via an -NO-dependent pathway (15). In this study, using a defined in vivo model of prefibrotic, asbestos-induced injury in the rat (7, 16), we show for the first time that inhalation of asbestos fibers stimulates the production of RNS in the lungs and pleura. Furthermore, we demonstrate that chrysotile as well as crocidolite asbestos fibers can induce the formation of nitrotyrosine, a marker of ONOO⁻ produc-

1. Abbreviations used in this paper: BAL, bronchoalveolar lavage; iNOS, inducible nitric oxide synthase; -NO, nitric oxide; O₂⁻, superoxide anion; -OH, hydroxyl radical; ONOO⁻, peroxynitrite anion; RNS, reactive nitrogen species; TNM, tetranitromethane.

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tion, at sites of asbestos fiber deposition as well as in the airway epithelium and pleural mesothelium.

Methods

Mineral dust samples and reagents. National Institute of Environmental Health Sciences (NIEHS, Research Triangle Park, NC) crocidolite fibers were a generous gift from Dr. Roy Rando (Tulane University School of Public Health, New Orleans, LA). Chrysotile fibers were obtained from the NIEHS. 3-nitro-1-tirosine, 3-aminoo-tirosine, O-phospho-l-tirosine, tetraniromethane (TNN), bovine serum albumin fraction V (BSA), sodium deoxycholate, phenylmethylsulfonyl fluoride, aprotinin, pepstatin A, sodium orthovanadate (NaVO₃), NaCl, EGTA, NaF, Na₂CO₃, NaHCO₃, KH₂PO₄, and Harris’s hematoxyl were purchased from Sigma Chemical Co. (St. Louis, MO). Tris base, Triton X-100, blocking reagent, and leupeptin were supplied by Boehringer Mannheim (Indianapolis, IN). Tween 20 was bought from Bio-Rad Laboratories (Herces, CA). Diff-Quik® Stain Set was obtained from Baxter Healthcare (McGaw Park, IL), and sodium pentobarbital was purchased from Abbott Laboratories (North Chicago, IL). RPMI 1640, FBS, and PBS were supplied by Biofluids, Inc. (Rockville, MD).

Animals and inhalation exposure regimen. For the inhalation exposures, three groups of male Fischer-344 rats were placed in whole-body inhalation chambers and exposed to either NIEHS crocidolite asbestos fibers, NIEHS chrysotile asbestos fibers, or filtered room air ( sham-exposed group). The rats in each group were comparably matched for age (56 ± 20 g) and size (mean weight for each group: 207 g) before inhalation exposure. Using an established inhalation exposure protocol (7, 16, 17), each group was exposed for 6 h/d over 2 wk. The rats were killed at 1 or 6 wk after the cessation of exposure by the intraperitoneal administration of sodium pentobarbital (50 mg/kg), followed by exsanguination via the abdominal aorta.

Total fiber mass aerosol concentrations were measured on membrane filters using 25-mm conductive cowl filter holders (Nuclepore, Carmel, CA) equipped with 0.8-µm pore filters (DM-800; Gelman Sciences, Ann Arbor, MI) by standard gravimetric analysis. All total mass measurements were on a daily basis at the nose level of the exposed animals. The asbestos aerosols were generated using a modified Timbrell dust generator (BGI, Waltham, MA). Aerodynamic particle-size distributions were measured using a Sierra 8-stage cascade impactor (model 210; Andersen Instruments, Atlanta, GA) throughout the exposure period. For one series of experiments, in which the rats were killed 1 wk after the cessation of exposure, the time-weighted exposure concentrations were as follows (mean ± SE): crocidolite 6.88±1.77 mg/M³; chrysotile 8.25±1.54 mg/M³. For another series of studies, in which the rats were killed 6 wk after the cessation of exposure, the time-weighted exposure concentrations were as follows: crocidolite 7.65±1.08 mg/M³; chrysotile 9.24±1.55 mg/M³. These asbestos exposure levels were comparable with historic asbestos dust concentrations recorded in the workplace environment of asbestos mines and mills (18). Analysis of all impactor data indicated the following: crocidolite geometric mean diameter = 0.4 µm, geometric SD = 3.8; chrysotile geometric mean diameter = 0.5 µm, geometric SD = 4.1. Thus, the aerosols were very similar in aerodynamic size distributions and were highly respirable.

Collection of pleural and bronchoalveolar lavage (BAL) cells. BAL and pleural lavages were performed on rats killed 1 wk after the termination of inhalation exposure. Lavages performed on the rats killed at 6 wk after exposure had been used previously for another study (17). Consequently, immunohistochemical studies could only be performed on the lungs but not on the lavage cells from those three groups of exposed animals in the current study.

The technique of pleural lavage was performed, as described previously (17). Briefly, an 18-gauge Teflon cannula was inserted into the pleural cavity via a small diaphragmatic aperture and each pleural space was lavaged twice with 5 ml of PBS, prewarmed to 37°C. BAL was carried out by inserting a tube into the trachea, whereupon the lung was lavaged six times with 7 ml of PBS, prewarmed to 37°C, as described elsewhere (19). Cells were pelleted by centrifugation at 500 g and then resuspended in PBS for evaluation of viability (by trypan blue dye exclusion) and yield (by hemocytometry).

Cytosin preparations of BAL cells, comprising >90% macrophages, were obtained using a Cytospin® 3 cytocentrifuge (Shandon, Pittsburgh, PA) and were processed for differential cell counts (with Diff-Quik® stain) and for immunocytochemical studies for inducible nitric oxide synthase (iNOS) protein expression. Enriched populations of pleural macrophages were obtained by adding crude pleural lavage cell populations (3 × 10⁶ cells/well) to 4-well chamber slides (Lab-Tek, Naperville, IL) containing RPMI 1640 + 10% FBS, and incubating them in 5% CO₂ at 37°C for 2 h. Thereafter, the cells were washed vigorously three times with PBS to remove nonadherent cells. Then the adherent cells, consisting of at least 77% macrophages, were used for immunocytochemical studies.

Measurement of NO₂ production by BAL cells. Washed BAL cells were obtained from rats that were killed 1 wk after the termination of the inhalation exposure regimen. BAL cells from two to three rats per exposure group were pooled, suspended in DMEM + 10% FBS, and added to 24-well tissue culture plates (3 × 10⁶ cells/well). After allowing the cells to attach for 1 h at 37°C, nonadherent cells were removed, 1 ml of DMEM was added, and the adherent cells were incubated for an additional 24 h at 37°C. Conditioned medium samples were collected and analyzed for NO₂ content, the oxidation product of NO, by the Griess reaction (20).

Processing of tissues for histology and immunohistochemistry. The lungs, diaphragms, and chest walls were excised from four rats in each exposure group that were killed at 1 or 6 wk after the cessation of the exposure regime. None of these animals had been subjected to BAL at the time of killing. The diaphragm and chest wall samples were fixed in 10% neutral buffered formalin. The lungs were inflation-fixed in situ by the intratracheal instillation of ~5 ml of 10% neutral buffered formalin at a pressure of 20 cm. Sagittal sections then were prepared of the left lung with sharp razor blades. All tissue samples were embedded in paraffin and routinely processed for histologic and immunohistochemical analyses, for which 5-µm-thick sections were prepared. Lung histology was evaluated on sections stained with hematoxylin and eosin and with the trichrome stain. Immunostaining for ED1 antigen, iNOS protein, and nitroty-rosine. Immunostaining was performed on histologic sections and on lavage cell preparations. Before performing immunocytochemical studies, the BAL cells on cytosin slides and the pleural lavage adherent cells on 4-well chamber slides were fixed with acetone for 10 min at ambient temperature. To inhibit endogenous peroxidase activity, the lavage cells (which had strong endogenous activity) were incu-bated with 3% H₂O₂ in absolute methanol, whereas the tissue sections (which manifested lesser endogenous activity) were incubated with 0.3% H₂O₂ in methanol for 30 min at ambient temperature. Specific immunocytochemical studies were performed using a monoclonal anti-iNOS protein antibody (1:20 dilution; Transduction Laboratories, Lexington, KY), a monoclonal anti-rat mononuclear phagocyte (ED1) antibody (1:50 dilution; PharMingen, San Diego, CA), and a polyclonal antinitrotyrosine antibody (1:50 dilution; Upstate Biotechnology, Lake Placid, NY) as primary antibodies for 1 h at ambient temperature. Since the anti-iNOS protein and anti-ED1 antibodies were not suitable for use in immunohistochemical studies on formalin-fixed sections, they were used exclusively for studies on BAL and pleural lavage cells. However, the antinitrotyrosine antibody was used on tissue sections because nitrotyrosine immunoreactiv-ity can be readily demonstrated in formalin-fixed tissue (21). The cell preparations and sections were labeled using a Universal DAKO LSBA® 2 kit containing an appropriate, biotinylated secondary antibody and streptavidin-conjugated peroxidase (DAKO, Carpinteria, CA). For immunocytochemical visualization, 3,3’-diaminobenzidine (Sigma Chemical Co.) was used as a substrate, and the cell preparations and sections were counterstained with Harris’s hematoxylin. As
a negative control, the appropriate specific primary antibody was omitted, and 1% blocking reagent was substituted. To confirm the specificity of the antinitrotyrosine antibody, the antiserum was incubated with either 10 mM 3-nitro-l-tyrosine, 10 mM 3-aminol-tyrosine, or 10 mM O-phospho-l-tyrosine in 0.05 M Tris-HCl buffer immediately before the addition of the primary antibody to the tissue sections. As an additional negative control, tissue sections were incubated with 10% normal rabbit serum in Tris buffer instead of the antinitrotyrosine antibody.

Quantitation of immunostained macrophages. Quantitative immunostaining for ED1 antigen, a specific phenotypic marker for rat mononuclear cells (22), and for iNOS protein expression was performed by counting the fraction of immunoreactive BAL and adherent pleural cells in 10 randomized microscopic fields at a magnification of 400. Since alveolar macrophages were readily identified in histologic sections, macrophage quantitation for nitrotyrosine immunoreactivity was performed by counting 400 consecutive alveolar macrophages in tissue sections at a magnification of 400 and then determining the proportion of immunoreactive cells.

Quantitation of lung nitrotyrosine content. Rat lung nitrotyrosine content was quantified in three rats from each exposure group that had been killed at 1 wk after exposure had ceased and that had not been subjected to BAL. For this purpose, the entire left lung was removed, placed in PBS on ice, and ground into a fine mesh using a vortex-style tissue mixer. After centrifugation at 3,000 g for 5 min at 4°C and removal of the supernatant, the tissue fragments were dispersed in five volumes of ice-cold tissue lysis buffer comprising the following: 50 mM Tris-HCl, pH 7.4, 1% Triton X-100, 0.25% sodium deoxycholate, 150 mM NaCl, 1 mM EGTA, 1 mM phenylmethylsulfonyl fluoride, 1 μg/ml aprotinin, 1 μg/ml leupeptin, 1 μg/ml pepstatin, 1 mM Na2VO4, and 1 mM NaF. The samples were sheared by sonication and stored overnight on ice. After centrifugation at 10,000 g for 10 min at 4°C, the supernatant was collected and assayed for lung protein and nitrotyrosine content. The protein concentrations of the lung sample supernatants were determined using a Bio-Rad Protein Assay Kit I (Bio-Rad Laboratories, Hercules, CA).

Quantitation of lung nitrotyrosine levels was performed by a modification of a previously published ELISA procedure (23). A nitratetreated protein solution was prepared for use as a standard by incubating 0.1% (1 mg/ml) BSA in 50 mM KH2PO4 at a pH of 7.4 for 30 min at 37°C with 1 mM TNM, a very efficient nitrating agent (24). After adjusting the pH to 10.0 with 3 M NaOH, the amount of nitrotyrosine present in the TNM-treated BSA solution was measured at 430 nm ($\epsilon_\text{e} = 4,400 \text{ M}^{-1}\text{cm}^{-1}$) and expressed as nanomoles of nitrotyrosine per milligram of BSA. Subsequently, a stock solution of the TNM-treated BSA was diluted with 0.1 M Na2CO3-NaHCO3, coating buffer at a pH of 9.6. These standard samples, along with tissue samples from the lungs of sham-exposed and asbestos-exposed rats (100 μl; 100 μg/ml), were applied to Immulon 2 ELISA plates (Dynatech, Chantilly, VA) and allowed to bind for 1 h at 37°C. After nonspecific binding sites were blocked with 1% BSA in PBS, the wells were incubated for 90 min at 37°C with a mouse IgG monoclonal antinitrotyrosine primary antibody (Upstate Biotechnology), diluted 1:200, and, subsequently, for 30 min at 37°C with a peroxidase-conjugated goat anti-mouse IgG secondary antibody (Kirkegaard & Perry Laboratories, Inc., Gaithersburg, MD), diluted 1:1,000. After washing the plates, the peroxidase reaction product was generated using a TMB Microwell Peroxidase Substrate (Kirkegaard & Perry Laboratories, Inc.). The reaction plate was incubated for 10 min at ambient temperature and its optical density was measured at 650 nm. We calculated the amount of nitrotyrosine content per milligram of protein by relating the optical absorbance values of the nitrated BSA standards measured by ELISA to the levels of nitrotyrosine per milligram of BSA as determined spectrophotometrically. Nitrotyrosine content of lung samples was expressed as picomoles of nitrotyrosine per milligram of protein.

Statistical analysis. Quantitative data are expressed as mean±SE of three experiments per group. The statistical significance of the data was determined by the Student’s t test.

Results

Total yield and composition of lavage cells. Table I shows the total cell yield, viability, and macrophage fraction obtained by BAL and pleural lavage from rats in each exposure group at 1 wk after the termination of exposure. Although there were no significant differences between the number of BAL cells recovered from rats in the three exposed groups, significantly greater numbers of total pleural cells and pleural macrophages were recovered from asbestos-exposed rats than from sham-exposed animals.

iNOS protein expression in BAL and pleural lavage cells. As shown in Fig. 1, at 1 wk after the cessation of exposure significantly higher percentages of iNOS-positive BAL cells were noted after both types of asbestos exposure than after sham exposure. More than 90% of these cells were macrophages and significantly greater total numbers of iNOS-positive alveolar macrophages were recovered from crocidolite- (1.51±0.13×106 cells/rat; $P < 0.05$ vs. sham-exposed) or chrysotile-exposed rats (1.89±0.11×106 cells/rat; $P < 0.005$ vs. sham-exposed) than from sham-exposed rats (1.02±0.13×106 cells/rat). Similar observations were noted in pleural lavage cells (Fig. 2) which comprised ~77–80% macrophages (Table II).

$\text{NO}_2^-$ production by BAL cells. The amounts of NO2 produced in 24-h cultures of BAL cells obtained from the three exposed groups of rats at 1 wk after exposure are shown in Fig. 1. Distinct differences between the groups were ob-

Table I. Total Cell Yield and Total Macrophages Recovered by BAL and Pleural Lavage from Asbestos-exposed and Sham-exposed Rats

<table>
<thead>
<tr>
<th>Exposure group</th>
<th>BAL cell viability and yield/rat ($\times 10^6$)*</th>
<th>Pleural lavage cell viability and yield/rat ($\times 10^6$)†</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Viability (%)</td>
<td>Total cells$^\text{a}$</td>
</tr>
<tr>
<td>Sham</td>
<td>89±1.3</td>
<td>6.3±0.83</td>
</tr>
<tr>
<td>Crocidolite</td>
<td>90±2.2</td>
<td>5.65±0.47</td>
</tr>
<tr>
<td>Chrysotile</td>
<td>94±0.4</td>
<td>6.85±0.41</td>
</tr>
</tbody>
</table>

* $n = 3$ rats/group. $^\text{a}n = 10$ rats/group. $^\text{†}$Mean±SE. $^\text{P} < 0.001$ compared with sham-exposed group. $^\text{§}$P < 0.01 compared with sham-exposed group. $^\text{**}$P < 0.05 compared with sham-exposed group.
served, as reflected by significantly greater NO$_2$ formation in BAL cell cultures from asbestos-exposed rats than from sham-exposed animals. Notably, chrysotile inhalation also was associated with significantly greater NO$_2$ production than was evident after crocidolite exposure ($P < 0.001$).

**Nitrotyrosine expression in lungs.** Examination of hematoxylin and eosin–stained sections of the lungs, chest wall, and diaphragms of sham-exposed rats revealed no histologic abnormalities. In contrast, the lungs of crocidolite- and chrysotile-exposed rats demonstrated prominent thickening of alveolar duct bifurcations with focal aggregations of macrophages at these sites, as has been described previously after asbestos inhalation (25, 26). These histologic findings were observed in lung sections obtained at both 1 and 6 wk after the cessation of asbestos exposure. However, no evidence of parenchymal, visceral pleural, or parietal pleural fibrosis was detected in the lungs of any of the exposed rats on Masson’s trichrome–stained sections.

Lung sections from asbestos-exposed rats also displayed strong immunoreactivity for the presence of nitrotyrosine at both 1 and 6 wk after the termination of exposure. Thus, diffusely positive cytoplasmic immunostaining for nitrotyrosine was evident in the region of alveolar duct bifurcations, both within alveolar duct epithelium and within underlying interstitial cells (Fig. 3). These findings were apparent after exposure to crocidolite (Fig. 3, A and B) as well as after chrysotile exposure (Fig. 3, C–E). Strong immunoreactivity for nitrotyrosine also was observed within the apical portions of the cyto-

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**Figure 1.** Evidence of iNOS activation in alveolar macrophages from sham-exposed and asbestos-exposed rats. Left ordinate shows the proportions of alveolar macrophages demonstrating iNOS immunoreactivity ($n = 3$). Right ordinate shows the production of NO$_2$ by cultured alveolar macrophages ($n = 10$). Values represent mean $\pm$ SE. *$P < 0.05$ compared with sham-exposed rats; **$P < 0.01$ compared with sham-exposed rats.

**Figure 2.** Immunoperoxidase staining for iNOS protein in pleural macrophages from a chrysotile-exposed rat at 1 wk after the cessation of exposure. Original magnification, $\times 150$.

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**Table II. Proportions of ED1-positive Cells and Total iNOS-positive Macrophage Yield in Pleural Lavages from Asbestos-exposed and Sham-exposed Rats**

<table>
<thead>
<tr>
<th>Exposure group</th>
<th>Pleural lavage cells$^a$</th>
<th>iNOS-positive cells$^b$</th>
<th>No. of iNOS-positive macrophages/rat ($\times 10^5$)$^c$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sham</td>
<td>79.5$\pm$0.8</td>
<td>4.41$\pm$0.55</td>
<td>3.49$\pm$0.15</td>
</tr>
<tr>
<td>Crocidolite</td>
<td>77.8$\pm$0.8</td>
<td>8.56$\pm$0.15$^d$</td>
<td>9.27$\pm$0.91$^d$</td>
</tr>
<tr>
<td>Chrysotile</td>
<td>78.8$\pm$0.7</td>
<td>9.59$\pm$0.11$^d$</td>
<td>8.66$\pm$0.87$^d$</td>
</tr>
</tbody>
</table>

$^a$Immunocytochemistry performed after nonadherent cells were eliminated. $^b$n = 10 rats/group. $^c$Mean$\pm$SE. $^dP < 0.0005$ compared with sham-exposed group. $^eP < 0.0001$ compared with sham-exposed group.
Figure 3. Immunoperoxidase staining for nitrotyrosine in lung sections from asbestos-exposed and sham-exposed rats at 1 wk after the cessation of exposure. (A) The bronchiolar epithelium (arrows) and alveolar duct region (arrowheads) of a crocidolite-exposed rat show positive immunostaining for nitrotyrosine. Original magnification, ×50. (B) Immunoreactivity for nitrotyrosine is observed in some of the interstitial cells and alveolar ductal epithelium of a crocidolite-exposed rat. Original magnification, ×150. (C) The bronchiolar epithelium and alveolar duct bifurcation region from a chrysotile-exposed rat show positive immunoreactivity for nitrotyrosine. Original magnification, ×50. (D) Positive immunostaining for nitrotyrosine is evident in many of the interstitial cells and in the alveolar ductal epithelium of a chrysotile-exposed rat.
plasm of the bronchiolar epithelium in both crocidolite-exposed (Fig. 3A) and chrysotile-exposed rats (Figs. 3C, and see Fig. 6A). In contrast, only faint immunoreactivity for nitrotyrosine was detected within occasional bronchiolar epithelial cells of sham-exposed rats at 1 wk (Fig. 3F) and 6 wk after exposure. A small proportion of alveolar macrophages in lung sections from sham-exposed animals displayed cytoplasmic immunostaining for nitrotyrosine (Fig. 4). However, the proportions of nitrotyrosine-positive alveolar macrophages detected in rat lung sections increased approximately four- to ninefold after both types of asbestos inhalation (Fig. 4). Cytoplasmic immunostaining of the alveolar macrophages in the lungs of exposed animals was intense (Figs. 3–5) and was seen at 1 wk and for as long as 6 wk after exposure had ceased (Fig. 4). Both crocidolite- and chrysotile-exposed rats also manifested focal areas of strong cytoplasmic immunoreactivity for nitrotyrosine in the visceral mesothelium (Fig. 5, A–C) as well as in the parietal pleural mesothelial cells lining the diaphragm (Fig. 5, D) and chest wall. Again, immunostaining for nitrotyrosine was evident within pleural mesothelial cells at 1 wk (Fig. 5, A and B) and at 6 wk (Fig. 5C) after the termination of exposure. No mesothelial immunoreactivity for nitrotyrosine was detected in tissue sections from sham-exposed rats.

Immunospecificity of the primary antibody for nitrotyrosine was confirmed when immunoreactivity of the tissues was abolished by absorption of antinitrotyrosine activity with soluble 10 mM nitrotyrosine (Fig. 6C). In contrast, neither 10 mM aminotyrosine nor 10 mM phosphotyrosine inhibited the binding of the antinitrotyrosine antibody (data not shown). This is in agreement with what has been described previously (11, 27). Furthermore, no tissue immunoreactivity was detected when 10% normal rabbit serum was substituted as the primary antibody (Fig. 6B).

Quantitation of lung nitrotyrosine content by ELISA. The total protein content of samples extracted from the left lung lobes of rats in each exposure group ranged from 2.11 to 2.23 mg. Significantly higher concentrations of nitrotyrosine were detected in the lungs of crocidolite- and chrysotile-exposed rats than in sham-exposed animals (Table III).

Table III. Lung Nitrotyrosine Content Determined by ELISA in Left Lung Lobes from Asbestos-exposed and Sham-exposed Rats at 1 wk after the Cessation of Exposure

<table>
<thead>
<tr>
<th>Exposure group</th>
<th>Lung nitrotyrosine concentration (pmol/mg protein)</th>
</tr>
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<tbody>
<tr>
<td>Sham-exposed</td>
<td>31±3.6</td>
</tr>
<tr>
<td>Crocidolite-exposed</td>
<td>59±5†</td>
</tr>
<tr>
<td>Chrysotile-exposed</td>
<td>84±12‡</td>
</tr>
</tbody>
</table>

* n = 3 rats/group. †Mean±SE. ‡P < 0.05 compared with sham-exposed group. §P < 0.01 compared with sham-exposed group.

Discussion

Our data show that inhalation of either chrysotile or crocidolite asbestos fibers results in increased levels of stable by-products of NO in BAL cells and the formation of nitrotyrosine within the lungs and pleura. Thus, 1 wk after the cessation of exposure, alveolar macrophage iNOS protein immunoreactivity and NO2 production were significantly greater after asbestos inhalation than after sham exposure. Notably, NO2 production in BAL cells was significantly greater after chrysotile inhalation than after crocidolite exposure, a finding that is consistent with a similar observation noted previously in rat pleural lavage cells (17).

Both groups of asbestos-exposed rats also manifested positive cytoplasmic immunostaining for nitrotyrosine in the region of thickened alveolar duct bifurcations as well as within bronchiolar epithelial cells, alveolar macrophages, and visceral pleural mesothelial cells. These findings are noteworthy, since the alveolar duct bifurcations are the sites of deposition of inhaled asbestos fibers and represent the locations where the earliest tissue response to inhaled asbestos fibers occurs (1, 25). In addition, nitrotyrosine immunoreactivity was observed within the cytoplasm of mesothelial cells lining the parietal pleura of the chest wall and the diaphragm. The parietal pleural location of the nitrotyrosine-positive foci in asbestos-exposed animals possibly may have pathogenetic significance, since asbestos-related parietal pleural plaques are detected in similar anatomic locations in circumstances of human occupational exposure to asbestos (1, 28). In contrast, the lungs of
sham-exposed rats manifested only faint, focal cytoplasmic immunoreactivity within bronchiolar epithelial cells and within occasional alveolar macrophages. Another important observation was the fact that nitrotyrosine immunoreactivity within the lung parenchyma and pleura was persistent, having been detected as soon as 1 wk and for as long as 6 wk after asbestos exposure had ceased.

Previous studies have demonstrated that nitration reactions may occur in the lungs after a variety of in vivo toxic insults such as in patients with adult respiratory distress syndrome (11) and in rats exposed to endotoxin (29), zymosan (12), or hyperoxia (11). These reactions have been attributed to the interaction of · NO with $O_2^-$ at a rate constant of $6.7 \times 10^9$ M$^{-1}$s$^{-1}$ at pH 7.4 to form ONOO$^-$ (30) and have been observed in circumstances associated with cytokine-induced inflammation. There is evidence that the nitration yield may be increased via the formation of the nitrosoperoxycarbonate anion ($O = N$-$OOCO_2^-$) from physiologic concentrations of CO$_2$ and bicarbonate, thereby enhancing the reactivity of ONOO$^-$ (31).

Significant pulmonary sources of · NO/ONOO$^-$ include activated macrophages (32), neutrophils (33), type II pneumocytes (34), and airway cells (34). It is now well-recognized that upregulated iNOS activity and · NO/ONOO$^-$ formation have been detected in rats (12) as well as patients with inflammatory lung disease (35, 36). Thus, it is pertinent that asbestos exposure in clinical settings as well as experimental asbestos inhalation models in the rat are characterized by protracted macrophage accumulations at sites of asbestos fiber deposition (1, 26, 28) and upregulated secretion of the cytokines, TNF-$\alpha$, IL-$\beta$, and interferon-$\gamma$, which induce iNOS activation (37–39). Moreover, chrysotile and amosite inhalation can induce proliferative lesions in rat airway and proximal alveolar epithelium (40, 41). Additionally, both chrysotile and crocidolite

Figure 5. Immunoperoxidase staining for nitrotyrosine in sections from asbestos-exposed rats at 1 wk (A, B, and D) and at 6 wk (C) after the cessation of exposure. (A) Immunoreactivity for nitrotyrosine is detected within the visceral pleural mesothelium and an occasional alveolar macrophage of a crocidolite-exposed rat. Original magnification, $\times$150. (B) Similar immunoreactivity for nitrotyrosine is evident within the visceral pleural mesothelium of a chrysotile-exposed rat. Original magnification, $\times$150. (C) Strong immunostaining for nitrotyrosine is observed in the visceral pleural mesothelium of a chrysotile-exposed rat. Original magnification, $\times$150. (D) Strong immunostaining for nitrotyrosine is observed in the parietal pleural mesothelium of a chrysotile-exposed rat at 1 wk after the cessation of exposure. Original magnification, $\times$150.
inhalation in the rat have been shown to induce a persistent macrophage inflammatory response within the pleural space associated with enhanced pleural macrophage TNF-α secretion (17). Therefore, the findings in the current study of iNOS activation and nitrotyrosine formation are likely to reflect the consequences of asbestos-induced pleuro-pulmonary cytokine activation.

At first glance, our levels of nitrated tyrosines in asbestos-exposed rats are small. However, it should be noted that these results are normalized to total lung protein whereas our immunocytochemical data show extensive nitration mainly in airway and mesothelial cells. Levels of lung nitrotyrosine detected in our studies are comparable with those reported by Leeuwenburgh et al. in human plasma-derived low-density lipoprotein (42), by Shigenaga et al. in normal rat serum (43), and by Khan et al. in human serum albumin and ONOO⁻-treated low-density lipoprotein in plasma (44). However, these concentrations are ~ 1–2 orders of magnitude less than those measured in atherosclerotic plaques (42) and the peritoneal exudate of zymosan-injected rats (43).

Although it is not possible to determine whether our observations are causally related to the induction of asbestos-related fibrosis and carcinogenesis, there is strong circumstantial support for this contention. Firstly, both chrysotile and crocidolite fibers were shown recently to induce the formation of ONOO⁻ in IL-1β-containing rat pleural mesothelial cell cultures, an effect that showed a good correlation with the cytotoxic action of those fibers on the cultured cells (14). The detection of protracted elevations of LDH levels in the BAL fluids of crocidolite- and chrysotile-exposed rats is consistent with these observations (19). ONOO⁻ is a potent oxidant that can initiate cell injury via oxidation of protein sulfhydryl moieties and lipid peroxidation (30) and by decreasing oxygen consumption and sodium uptake through amiloride-sensitive sodium channels (45, 46). Recently, it also has been demonstrated that ONOO⁻ can activate the enzyme poly (ADP-ribose) polymerase in human pulmonary epithelial cells, thereby inducing cellular energy depletion, inhibition of mitochondrial respiration, and increased cellular permeability (47). Notably, both chrysotile and crocidolite fibers were shown to induce poly (ADP-ribose) polymerase activation in rat pleural mesothelial cells (48). Additionally, there is evidence that ONOO⁻ can mediate apoptosis in diverse inflammatory settings (49, 50). This may have significance for asbestos-related injury, since two recent studies have documented that asbestos fibers can induce apoptosis in pleural mesothelial cells (51, 52) and, in one of them, apoptosis was abrogated by a poly (ADP-ribose) polymerase inhibitor (52).

Several studies have shown that ONOO⁻-induced protein nitration may compromise protein function and mediate tissue injury. In one, nitration of tyrosine residues of human IgG abrogated their C1q-binding activity (53). Tyrosine nitration also has been shown to inhibit protein phosphorylation by tyrosine kinases, thus interfering with signal transduction mechanisms (54), and to inhibit the assembly of neurofilament subunits by nitrating a major structural protein of motor neurons (55). Whatever mechanisms may be involved, there are strong parallels between the findings in our rodent model of asbestosis and those of Saleh et al. in patients with idiopathic pulmonary fibrosis (35). These authors reported the existence of signifi-
cant amounts of nitrotyrosine and iNOS in macrophages, neutrophils, and alveolar epithelial cells of patients with idiopathic pulmonary fibrosis and proposed that RNS may be responsible for the generation of fibrosis. The exact mechanisms whereby this occurs have not been identified.

It is conceivable that activation of iNOS within alveolar and pleural macrophages may be linked to the fibroproliferative and neoplastic effects of asbestos. In this regard, it is pertinent that ·NO has been shown to upregulate the release of IL-1 and TNF-α from mouse peritoneal macrophages (56). Also, by mobilizing iron from crocidolite fibers or from asbestos bodies, ·NO can produce DNA strand breaks (10) and DNA base modification (15). Moreover, rat pleural mesothelial cells stimulated with cytokines and LPS were found to generate potentially carcinoagenic N-nitrosating compounds from ·NO (57). Therefore, the detection in the present study of strong immunostaining for nitrotyrosine within the airway epithelium and visceral and parietal pleural mesothelium of asbestos-exposed rats may have added significance, since these epithelial cells are the progenitor cells for bronchogenic carcinoma and malignant pleural mesothelioma, respectively.

In summary, this study has demonstrated for the first time that inhaled asbestos fibers, which are recognized fibrogenic and carcinogenic agents, can induce the formation of RNS in the rat pleura and lung parenchyma. It is also noteworthy that the biologic effects of chrysotile were found to be equivalent to or greater than those of crocidolite asbestos in this regard. Therefore, our findings underscore the pathogenic potential of chrysotile to inflict pleuro-parenchymal injury (4, 5). Furthermore, the induction of RNS by both chrysotile and crocidolite fibers in vivo may provide an alternative mechanism of asbestos-induced injury to that thought to be induced by Fenton reactions.

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