In Situ Microscopic Analysis of Asbestos and Synthetic Vitreous Fibers Retained in Hamster Lungs following Inhalation

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Hamsters breathed, nose-only, for 13 weeks, 5 days/week, 6 hr/day, either man-made vitreous fiber (MMVF)10a, MMVF33, or long amosite asbestos at approximately 300 World Health Organization (WHO) fibers/cc or long amosite at 25 WHO fibers/cc. [World Health Organization fibers are longer than 5 μm and thicker than 3 μm, with aspect ratio >3.1] After sacrifice, fiber burden was estimated (left lungs) by ashing and scanning electron microscopy (ashing/SEM) or (right middle lobes) by confocal laser scanning microscopy (CLSM) in situ. In situ CLSM also provided three-dimensional views of fibers retained, undisturbed, in lung tissue. Fibers of each type were lodged in alveoli and small airways, especially at airway bifurcations, and were seen fully or partly engulfed by alveolar macrophages. Amosite fibers penetrated into and through alveolar septa. Length densities of fibers in parenchyma (total length of fiber per unit volume of lung) were estimated stereologically from fiber transsections counted on two-dimensional optical sections and were 30.5, 25.3, 20.0, and 81.6 mm/mm3 for MMVF10a, MMVF33, and low- and high-dose amosite, respectively. Lengths of individual fibers were measured in three dimensions by tracking individual fibers through series of optical sections. Length distributions of amosite fibers aerosolized, but before inhalation versus after retention in the lung were similar, whether determined by ashing/SEM or in situ CLSM. In contrast, the fraction of short MMVF10a and MMVF33 fibers increased and the geometric mean fiber lengths of both MMVF's decreased by approximately 60% during retention. Most likely due to fiber deposition pattern and differences in sampling, fiber burdens [MMVF10a, MMVF33, and amosite (high dose; 269 WHO fibers/cc)] determined by ashing/SEM were 1.4, 1.5, and 3.5 times greater, respectively, than those calculated from in situ CLSM data. In situ CLSM is able to provide detailed information about the anatomic sites of fiber retention and also fiber lengths and burdens in good agreement with ashing/SEM results. Key words: aerosol, amosite asbestos, ashing, confocal microscopy, fiber burden, man-made fibers, MMVF, respirable, three-dimensional. Environ Health Perspect 107:367–375 (1999). | Online 26 March 1999 | http://ehpnet1.niehs.nih.gov/docs/1999/107p367-375rogers/abstract.html

The health effects of exposure to airborne asbestos and synthetic fibers have been extensively studied, as reviewed by DeVuyst et al. (1) and Hesterberg et al. (2). The respiratory system's response to fibers depends greatly on fiber composition, fiber length, the sites of deposition, the tissue compartments through which fibers move, and the rates at which they are cleared (3,4). Such health effects also depend on the amount of fiber deposition in the respiratory system and the resulting fiber burden. This has traditionally been determined by recovering fibers from lungs using thermal ashing techniques or chemical digestion by strong bases and oxidizing agents (5–7). Despite a considerable body of work, little is known about either the distribution of artificial fibers within the lungs after exposure or the local responses of lungs to fiber deposition. Indeed, it has been difficult to obtain such information because, although digestion methods permit determinations of numbers and dimensions of all fibers retained in the lung, neither fiber position within pulmonary structures nor any relationship to regional lung pathology can be recovered. Also, visualization methods available hitherto have required that the lung be finely sectioned physically to examine the fibers directly. The mechanical properties of the fibers themselves make them difficult to approach without disturbing or breaking them; not only are they needlelike (inhaled fibers frequently have aspect ratios >5:1 and may be over 20 μm in length), they also may be both rigid and fragile. Thus, if samples of lung tissue are embedded in paraﬃn and physically sectioned in preparation for conventional transmission light microscopy, the fibers may be cut, disturbed, or torn out of the section during the cutting process, as fibers would typically not lie entirely within physical sections that may be 6–8 μm in thickness. Transmission electron microscopy (TEM) has been used to image fibers within fixed lung tissue (8,9), but here, especially, the process of obtaining physical sections for TEM (60–70 nm thick) can be expected to cut apart every fiber encountered and to introduce artifacts (6). Scanning electron microscopy (SEM) permits intact fibers to be studied, especially following freeze-etching or dissection along bronchi; however, SEM images show primarily the surfaces of fibers, cells, and tissue that are closest to the observer. Confocal microscopy, however, makes it possible to peer tens of hundreds of microns below the surface of a translucent specimen and recover two-dimensional images from essentially any location within it, even revealing fibers within cells or tissues. Data volumes (three spatial dimensions plus intensity values) built up from such two-dimensional images obtained from increasing depths can be presented as stereoscopic (quasi-three-dimensional) pictures showing, for example, intact, undisrupted fibers penetrating alveolar walls. The present study's purpose was to determine the numbers and locations of three types of fibers [two man-made vitreous fibers (MMVF's) and long amosite asbestos, a naturally occurring mineral fiber] that had been inhaled into hamster lungs. We utilized confocal microscopy to examine relatively large embedded lung tissue samples with minimal disruption of the fibers deposited there and compared the results to those obtained by SEM of ashing residue.

Materials and Methods

Experimental Design

This article presents results obtained by examining material produced as part of a larger study. The parent study was an 83-week chronic inhalation study with interim sacrifices; the first one, after 13 weeks of exposure, supplied the material for the present investigation. There were 125 hamsters in each of five exposure groups and 140 in the exposure and control groups, respectively. On the 13th week, the animals were killed, and lungs were removed for examination. The experimental setup was identical to that used for the study of DeVuyst et al. (1), and the hamsters were housed in stainless steel cages to prevent contamination of the environment. Material was embedded in either paraffin or agar for histological sections or in paraffin or paraffin-free for TEM and SEM. TEM was used to observe the fibers within the tissue, whereas SEM was used for the surface morphology of the fibers. SEM images were obtained using an Hitachi S-570 microscope with a Hitachi TF-32000 electron gun. The samples were sputter-coated with a thin layer of gold-palladium before examination. The images were recorded on photographic film and digitized using an Image-Pro Plus computer program. The images were then analyzed using a Quantumahan program to determine the number and dimensions of the fibers. The results were then compared to those obtained by SEM of the ashing residue.
an air control group. Some results from the parent study have been published (10).

Experimental Material

Fibers. Three types of fibers were studied: 1) MMVF10a, a glass fiber commonly used in building insulation; 2) MMVF33, a glass fiber used in the manufacture of high-efficiency air-purification filter systems; and 3) long amosite, a form of naturally occurring mineral asbestos. World Health Organization (WHO) fibers have aspect ratio >3, length >5 µm, and diameter <3 µm (11).

Animals. Male Syrian golden hamsters (strain LakCr:LVG) of specific-pathogen-free quality were housed and exposed to aerosolized fiber at Research Consulting Co. (Füllinsdorf, Switzerland). They were housed individually in Makrolon type-1 cages (EHRET GmbH, Emmendingen, Germany) containing dust-free, softwood bedding (Lignocel, Schill AG, Muttenz, Switzerland) in a specific-pathogen-free environment. Pelleted food and water were provided ad libitum. This breed has been used in previous inhalation studies with natural and/or synthetic vitreous fibers (12).

Experimental Exposure

Animals were divided into six groups (five exposure groups and one air control group). Group 1 (MMVF10a fibers) was composed of animals exposed to 339 WHO fibers/cc (29.6 mg/mm³); group 2 (MMVF33 fibers) was composed of animals exposed to 310 WHO fibers/cc (37.0 mg/mm³); group 3 (long amosite, low concentration) animals received 36 WHO fibers/cc (0.8 mg/mm³); group 4 (long amosite, medium concentration) animals received 165 WHO fibers/cc (3.7 mg/mm³); group 5 (long amosite, high concentration) animals were exposed to 269 WHO fibers/cc (7.1 mg/mm³); and group 6 (air control) animals received no fibers, only air. For nose-only exposure to aerosols or filtered air, all animals were placed in restraint tubes for 6 hr/day, 5 days/week for 13 weeks; during the rest of the time, they breathed clean filtered room air. One day after the exposure period, hamsters were anesthetized with an intraperitoneal injection of sodium pentobarbital (300 mg/kg body weight) and sacrificed by exsanguination, whereupon their lungs were excised.

Specimen Preparation for in Situ Microscopic Examination

Fixation and embedding. The middle lobe of the right lung of each hamster was processed for microscopic examination of fibers in situ. The right lung was detached between lung and hilum, fixed by instilling 4% buffered paraformaldehyde into the airways at 30 cm water pressure, then shipped from Switzerland to Boston. The lung samples remained in the fixative for up to 10 days. Each right middle lobe was cut sagitally into five pieces 5 mm thick, dehydrated in graded ethanolic series to absolute ethanol, stained nonspecifically with the fluorescent dye lucifer yellow CH [0.001% w/v in absolute ethanol; (13)], and embedded in Spurr’s epoxy (Polysciences, Warrington, PA). The results of Law et al. (14) suggest that at most 16% of the silica content of the most susceptible MMVF3s would be dissolved in this time, and that the loss could be less than 1% from all fiber types.

Sample manipulation. Epoxy-embedded samples were prepared by a metal adapter plate and mounted on a high-speed milling apparatus, where the top of the block was milled to a glassy smoothness (15). The sample on its adapter plate was then attached to a stage with digital readouts on the confocal laser scanning microscope/microscopy (CLSM). The CLSM can only retrieve optical sections to a certain depth, the depth limit depending on objective working length and the severity of light scattering and absorption by the sample. Material below this depth limit could be imaged by returning the sample block to the milling apparatus to remove some of the already imaged overlying tissue. Upon its return to the CLSM, the block could be positioned in three dimensions to within 1 µm of its previous position.

Specimen Preparation for Ash Analysis

The left lung (detached between lung and hilum) was frozen at autopsy and shipped frozen from Switzerland to the Johns Manville International, Inc., analysis laboratories (Littleton, CO). To recover inhaled fibers from the lung for these evaluations, tissue from the left lung was subjected to a low-temperature ashing process as previously described (7,10,16); the ash residue was suspended in deionized filtered water, sonicated, and filtered onto a polycarbonate membrane filter (Nucleopore; Bio-Rad Laboratories, Hercules, CA: 0.2 µm pore size), which was mounted on a specimen plachet, coated with gold, and examined by SEM. Fibers were counted and measured according to methods described previously (10).

Imaging of Fibers and Lung Tissue by CLSM

CLSM configuration. The CLSM was a Sarastro 2000 (Molecular Dynamics, Sunnyvale, CA) fitted with a 25-mW argon ion laser providing 488 nm wavelength excitation illumination. Either 20× (NA 0.8) or 60× (NA 1.4) objectives were used; a polarizing filter placed in the emitted light’s path transmitted properly polarized light reflected from fibers.

The CLSM was configured with two channels, each with a separate photomultiplier detector. In channel 1, used for fluorescence (tissue) detection, light (wavelengths >510 nm) emitted from dye-stained tissue was directed to detector 1. In channel 2, used for reflected light (fiber) detection, illuminating light reflected from fibers (wavelengths <510 nm) passed through a 510-nm secondary beam splitter and was directed to detector 2.

Each field-of-view (FOV), therefore, was represented by two digital images—one from each channel—and such pairs of images could be superimposed to show fiber positions in relation to lung tissue. Optical sections lay in the microscope’s (x,y) focal plane.

Fiber samples: imaging by CLSM. Samples of dry stock MMVF10a, MMVF33, and long amosite asbestos fibers were obtained from the NAIMA (North American Insulation Manufacturers Association, Alexandria, VA) fiber repository and were suspended (1 mg/ml) in saline and allowed to settle in microwell chambers for CLSM. Micrographs of fibers were obtained with the CLSM configured as described previously, using the 20× and 60× objectives.

Lung tissue and fibers retained in lung: imaging by CLSM. The CLSM was used to look through the smooth surfaces of the epoxy-embedded tissue samples to as much as 150 µm below the surface without physically sectioning the samples.

Stacks of aligned optical sections were acquired between two depths sufficiently separated to completely contain essentially all fibers detected within them. This technique permitted us to determine the length densities, length distributions, and anatomical placement of retained long amosite and MMVF fibers in lungs. Three-dimensional projections were generated from stacks of optical sections using Voxel View software (Vital Images, Inc., Fairfield IA) (17). CLSM images for stereologic length densities. Length density refers to the length of a linear structure present within a unit volume of some containing phase; here, it refers to the total length of retained fiber per unit volume of lung and has units such as millimeters per cubic millimeter. Micrographs were obtained with the CLSM configured as described previously, using the 20× objective. Each digital micrograph was a two-dimensional, 1,024 × 1,024 array
520.0 μm on a side of gray-scale pixels (intensity values from 0 to 255), 0.5078 μm on a side; thus, pixel area was 0.2579 μm² and image area 270,399 μm². FOVs were acquired at random locations over the tissue exposed at the prepared block face.

CLSM images for three-dimensional lengths of individual fibers. Each image, acquired with the 60x objective, was a 512 × 512 array 87 μm on a side of gray-scale pixels, 0.34 μm on a side. The images were obtained with a vertical spacing of 0.7 μm. Stacks encompassing >75 μm along the z-axis were obtained from locations where fibers were relatively plentiful.

Image segmentation to emphasize fibers in the presence of tissue. Prior to analysis, each pixel in the field of view was assigned to one of three categories: tissue, fiber, or other. This was done by setting pixels representing air and noise in the channel 1 (tissue) image to 0, leaving tissue pixels with values >0. Fiber pixels in channel 1 were 0-valued. The channel 2 (fiber) image was similarly thresholded, leaving >0 values in pixels representing fiber; because of the slight reflectivity of the tissue and weak fluorescent emissions occurring at short wavelengths, some tissue appeared faintly in channel 2. Pixels representing fibers could be distinguished as those that were 0 in channel 1 but of at least moderate intensity in channel 2; side-by-side comparisons of the channel 1 and 2 images made it easy to identify fiber profiles in the channel 2 image.

Length Densities of Fibers within the Lung, by in Situ CLSM

Fiber length densities (total length of retained fiber per unit volume of lung) were measured in four, five, five, and five animals from the MMVF10a, MMVF33, low-dose amosite, and high-dose amosite treatment groups, respectively (13-week exposure; no recovery period). Images were taken predominantly of the two tissue samples with large cut surfaces on both sides and were collected at random (regular grid with randomly positioned origin); exception to this random procedure was made in that FOVs with visible pleura or extrapulmonary space were not recorded. Fiber transections visible on each image were counted manually.

Figure 1. Three types of fibers before aerosolization imaged by confocal laser scanning microscopy operated in reflected light mode. Individual fibers of all types appear to be of constant diameter. (A) Man-made vitreous fiber (MMVF)10a: fibers appear somewhat narrower and more curved than MMVF33. (B) MMVF33: note aggregates of individually straight fibers. (C) Long amosite: straight fibers, apparently little aggregated. Bars represent 10 μm.

Figure 2. Hamster lungs after 13-week exposure to three fiber types. Optical sections acquired approximately 50 μm below cut surface of embedded tissue using confocal laser scanning microscopy operated in dual detector mode (reflected light for fibers, represented by red in figure; fluorescence for tissue, represented by grayscale in figure). At least 500 μm of epoxy with embedded tissue was milled away from the surface (see text) prior to examination to ensure that fibers seen here could not have been disturbed by the act of cutting samples from the fixed lung before embedment. (A) Man-made vitreous fiber (MMVF)10a: 31 fiber transections seen in alveolar parenchyma; 21 are >5 μm from nearest septal profile. (B) MMVF33: 31 transections of fibers (arrowheads) are seen within alveolar parenchyma; of these, 21 are more than approximately 5 μm from the nearest alveolar septal profile. Lung pleura is visible. (C) Long amosite: seven fiber transections (arrowheads) and two alveolar macrophages (arrows) can be seen; one of the macrophages contains at least three fibers, the other, at least one. Bars represent 50 μm.
We assume that fibers retained in lung are randomly oriented with respect to the optical plane of section. The total length of linear features of interest (fibers) contained within a phase (lung) may be estimated \( (18) \) by

\[
L_V = 2 \times \frac{\sum_{i=1}^{M} Q_{fi}(i)}{\sum_{j=1}^{N} A_{lung}(j)} \tag{1}
\]

where \( L_V \) is the length density of fibers in lung, i.e., length of fibers per unit volume lung; \( Q_{fi}(i) \) is the number of fiber transections seen on image \( i \); and \( A_{lung}(j) \) is the transected area of lung seen on image \( j \) (here, always equal to the area of an image \( (18) \), Eq. 4.54a)).

**Fiber Dimensions**

**Fiber dimensions before aerosolization, as determined by SEM.** Lengths and diameters (analyzed using SEM and reported as mean, standard deviation, minimum, maximum, and median) of fibers to be aerosolized were obtained from NAIMA.

**Dimensions of retained fibers, as determined by ashing/SEM.** Numbers and dimensions of fibers after aerosolization, inhalation, retention, andashing of the left lung were determined by analysis of SEM micrographs of fiber samples collected on Nucleopore polycarbonate filters (0.2 \( \mu \)m pore size) \( (19) \).

**Fiber lengths as determined by CLSM in situ.** The lengths of fibers retained in animals’ right lungs were determined from fibers contained in stacks of optical sections. The length of a fiber was estimated as the sum of Pythagorean distances between transections on successive optical sections, whose locations were marked in three dimensions (pixel located at coordinates \( x,y \) on section at depth \( z \)).

**Fiber Burden**

**Fiber burden calculated from CLSM data.** The number of fibers in parenchyma per lung was estimated by combining estimates of fiber length density and geometric mean fiber length with an assumed lung volume. (Lung volume was not measured directly.)

\[
FB = \frac{L_V \times V_L}{\lambda_{fiber}} \tag{2}
\]

where \( FB \) is the fiber burden, number of fibers per left and right lung pair; \( L_V \) is the length density of fibers in lung, from two-dimensional CLSM data and stereology; \( V_L \) is the volume of lung, assumed to be 4.0 ml; \( \lambda_{fiber} \) is estimated geometric mean length of fibers, from three-dimensional CLSM data.

**Results**

**Characteristics of Fibers before Aerosolization**

Nonaerosolized samples of the three fiber types were examined by reflected light confocal microscopy. In Figure 1A (MMVF10a), the diameters along any given fiber appeared constant, but the fibers themselves occasionally were curved. Fibers occurred with a range of diameters, however. In Figure 1B (MMVF33), individual fibers appeared straight, often aggregated side to side and jumbled together. In Figure 1C (amosite), fibers appeared as straight rods of constant diameter, rarely clustered. All fiber types displayed brightness that varied depending on fiber orientation relative to the imaging plane, and they revealed interference patterns characteristic of fibers illuminated by polarized light. Practically the full lengths of the fibers were seen in these images, despite the optical sectioning property of confocal microscopy, because the fibers lay on a flat surface parallel to the optical sectioning plane. Because all fibers seen were in focus simultaneously, they must have been coplanar, and there was no indication of out-of-plane unobserved fibers. Hence, it is most likely that all fibers were clearly imaged and observed.

**Placement of Retained Fibers**

Samples of hamster lung following 13-week inhalation exposure to the three fiber types were examined using combined fluorescence and reflected light confocal microscopy. Fibers of each type reached the alveolar parenchyma and in some instances were taken up by alveolar macrophages (Fig. 2). Both MMVF10a (Fig. 2A) and MMVF33 (Fig. 2B) appeared throughout alveolar parenchyma, even occurring in subpleural regions. Fiber transections often were seen some tens of microns away from the nearest septal profile, meaning that the fiber did not lie against an alveolar septum for its entire length but crossed an alveolar airspace. With

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*Figure 3. Man-made vitreous fiber (MMVF10a) fibers retained for up to 13 weeks in hamster lung, visualized by confocal laser scanning microscopy. Bars in (A) and (B) represent 20 \( \mu \)m. (A) A single optical section from a stack of serial optical sections obtained at 1-\( \mu \)m intervals. Inflammatory cells with partially or wholly ingested fibers (in red) are in the airspace. (B) Left and right panels: stereo projection of the entire stack. Fibers are visible both in inflammatory cells and free in airspace.*
high-dose amosite (Fig. 2C), some fibers were found within alveolar macrophages (arrow, Fig. 2C).

Fibers of each type had reached alveolar parenchyma. Fiber placement within parenchyma was further established by examining stereoscopic views of stacked, aligned optical sections. For MMVF10a (Fig. 3A, 3B), fibers were present both free in the airspace and partially or wholly ingested by inflammatory cells. For MMVF33 (Fig. 4A, 4B), as with MMVF10a, some fibers were engulfed by inflammatory cells within alveolar air spaces (Fig. 4A, hollow arrow). A long fiber lay within the interstitium of an alveolar septum (Fig. 4A, arrow). The accompanying stereo projection (Fig. 4B) shows fiber position and orientation in alveoli. In amosite (Fig. 5A, 5B), a fiber (Fig. 5A, arrow) appears within the interstitial space adjacent to an airway. The stereo projection (Fig. 5B) shows this fiber to be entering the interstitium at an airway bifurcation (Fig. 5B, arrow). Numerous inflammatory cells containing fibers are present in the airway.

An alveolar macrophage can be seen attempting to engulf a long amosite fiber in Figure 6, which was acquired at higher magnification than Figures 3, 4, and 5. Most of the amosite fiber’s length is either embedded in sepal tissue or surrounded by the macrophage’s cell body or its slender tapering processes.

### Length Distributions of Fibers as Aerosolized and after up to Thirteen Weeks of Retention

Table 1 summarizes lengths of MMVF10a, MMVF33, and long amosite asbestos fibers as measured in air and lung samples 1) present in the aerosol presented to the animal’s nose; 2) after 13 weeks of exposure, by the ashing/SEM technique; and 3) after 13 weeks of exposure by in situ CLSM.

The order of aerosolized fiber lengths from longest to shortest (for both geometric mean lengths and median lengths) was MMVF10a, MMVF33, amosite; for retained fibers, the order was different—amosite, MMVF10a, MMVF33. Both MMVF types were shortened by approximately 60% (geometric mean). The geometric mean for amosite increased by 44%, whereas the arithmetic mean and median decreased by 9 and 6%, respectively. All fiber length distributions are skewed toward longer lengths (see below).

Figure 7 compares fiber length cumulative frequency distributions (FLCFDs) 1) for fibers in the aerosol, determined using SEM; 2) for retained fibers, using ashing/SEM; and 3) for retained fibers, using in situ CLSM. With MMVF10a (Fig. 7A), both analyses of retained fibers yielded similar LCFDs and both found fewer long fibers than did aerosol analysis; in particular, 37% of fibers in the aerosol were longer than 20 µm while barely 5% of retained fibers were. With MMVF33 (Fig. 7B), ashing/SEM and in situ CLSM did not yield similar LCFDs for retained fibers; the former showed a smaller proportion of short (<5 µm) fibers than did the latter. Both, however, showed fewer long fibers (>15 µm) than did aerosol analysis. With amosite asbestos (Fig. 7C), ashing/SEM analysis of retained fibers showed more short (<5 µm) and fewer long (>20 µm) fibers than did in situ CLSM. Neither analysis of retained fibers, however, showed as many long fibers as did aerosol analysis.

Figure 8 shows the complete LCFD obtained by in situ CLSM for retained fibers of each type. The fraction of total fiber number lying between any two lengths may be read directly from Figure 8. For example, the proportion of fibers categorizable as respirable according to WHO standards (identical with the category of all fibers longer than 5 µm and listed in Table 2) may be obtained as the fraction of theordinate above the crossing point of the LCFD and the vertical dotted line at 5.0 µm. Similarly, these curves demonstrate that the proportion of long (>20 µm) fibers was less than 5% for MMVF10a, nearly 15% for MMVF33, and 25% for amosite.

### Length Densities of Fibers in Lung by In Situ CLSM

Table 3 gives the number of FOVs examined, the number of fiber transections found, and the calculated length density (Eq. 1) for each fiber treatment group. MMVF10a and MMVF33 apparently
behaved similarly during retention, according to their distributions (given previously) and Table 3 demonstrates that length densities of these fibers were also comparable.

**Fiber Burden**

Fiber burdens (Eq. 2) of MMVF10a, MMVF33, and amosite (high dose) at the end of the 13-week exposure were estimated by two different techniques (ashing/SEM and in situ CLSM). Results are presented in Table 2, which lists the fraction of total fiber number in each 5-μm-wide length class up to 20 μm and the >20 μm length class; these numbers are the same as plotted in Figure 6 and were used to calculate the corresponding fiber burdens. The ratios between the fiber burdens for the two techniques are also provided for each fiber type. Discrepancies between the burdens are greatest for amosite. The burdens for amosite were two- to nine-fold greater with SEM than with CLSM, with the short fiber category (<5 μm long) showing the greatest disparity; in millions of fibers/lung, the burdens were 17 by SEM as compared to <2 by CLSM (Table 2). The two methods yielded fairly similar burdens for the two MMVFs, with SEM burden no more than 2.4-fold greater than CLSM. The distribution of the fiber burden by length class was similar for both MMVFs, except that MMVF33 had a greater number of long (>20 μm) fibers than did MMVF10a.

**Discussion**

**Locations of Fibers Retained in Lung, by in Situ CLSM**

We observed the placement of MMVF and asbestos fibers in situ in lung tissue after 13 weeks of exposure. Fibers of all three types

| Table 1. Fiber lengths before and after retention in lung |
|--------------------------------------------------------|
| Fiber type, experimental condition | Number of fibers measured | Arithmetic mean (SD) (μm) | Geometric mean (SD) (μm) | Mean (μm) | Range (μm) |
| MMVF10a | | | | | |
| Aerosolized (at nose, SEM)* | – | 19.4 (20.9) | 12.4 (0.916) | 12.3 | – |
| Postretention (ashing/SEM)* | – | 7.98 (6.18) | 6.4 (1.9) | 6.12 | – |
| Postretention (in situ CLSM) | 256 | 6.95 (6.20) | 5.15 (0.765) | 4.66 | 0.98–59.0 |
| Percent decrease | – | 64.2 | 58.5 | 68.8 | – |
| MMVF33 | | | | | |
| Aerosolized (at nose, SEM)* | – | 16.9 (16.3) | 11.8 (0.832) | 11.9 | – |
| Postretention (ashing/SEM)* | – | 9.97 (8.38) | 7.6 (2.1) | 7.15 | – |
| Postretention (in situ CLSM) | 250 | 9.40 (14.1) | 4.77 (1.12) | 3.50 | 0.51–74.8 |
| Percent decrease | – | 44.4 | 59.6 | 67.3 | – |
| Long amosite | | | | | |
| Aerosolized (at nose, SEM)* | – | 18.4 (16.7) | 8.6 (0.875) | 7.8 | – |
| Postretention (ashing/SEM)* | – | 11.68 (9.68) | 8.4 (2.1) | 8.88 | – |
| Postretention (in situ CLSM) | 396 | 18.8 (15.2) | 12.4 (0.775) | 12.08 | 2.67–113 |
| Percent decrease | – | 8.7 | 44.2 | 6.4 | – |

Abbreviations: SD, standard deviation; MMVF, man-made vitreous fiber; SEM, scanning electron microscopy; CLSM, confocal laser scanning microscopy. Descriptive statistics for lengths of three types of fibers (MMVF10a, MMVF33, and amosite asbestos) as present in aerosol administered to hamsters and from hamster lungs after 13 weeks of exposure, as determined by ashing/SEM or in situ CLSM.

*Results from Miller (19).
were deposited in small airways and alveolar septa, frequently being seen at airway bifurcations; this agrees with observations of Brody et al. (8) and Brody and Roe (20). Long amosite asbestos fibers frequently penetrated into alveolar septa as far as the interstitium and even completely through septa, in accord with the demonstrated ability of fibers to penetrate lung tissue (21–23). MMVF fibers were not observed to penetrate septa as frequently as did amosite fibers, however. Fibers of all types were found within alveolar macrophages.

Effects of Length-biased Filtering during Respiration and Fiber Fragmentation and Degradation during Retention

As in previous rodent inhalation studies (24), the present study demonstrated that both arithmetic and geometric mean lengths of MMVF fibers in aerosol were typically longer than those of retained fibers. This difference could be the result of a lung filtering process in which longer fibers are prevented from being inhaled into the distal alveolar parenchymal portions of the lung, or it could be a result of fiber degradation within the lung, or both.

In the case of amosite (Fig. 7C), the aerosol FLCFD and retained FLCFD obtained by in situ CLSM were similar, suggesting that amosite fibers of all lengths were equally likely to reach parenchyma—that is, that no significant filtering process affected amosite fibers. Because the FLCFD of retained amosite fibers was obtained after 13 weeks of exposure—adequate time for fiber fragmentation and degradation to have occurred in the lung—it seems that amosite fibers also resisted breakage and degradation and remained intact.

MMVF10a and MMVF33, on the other hand, showed strong reductions in arithmetic and geometric mean lengths (Table 1) and a strong increase (Fig. 7A, 7B) in the proportion of short fibers over the same time period. The 13-week time period cannot shed much light on which process (filtration or degradation) is more important in causing these changes because the possible filtration effects could be masked by effects of degradation. However, as part of the chronic study, fiber lengths were determined after only 6 hr of retention (10,19), and the proportion of long fibers by fiber type was as follows: MMVF10a: aerosol=6 hr > 13 weeks, suggesting that both lung filtration and fiber degradation were contributing factors; MMVF33: aerosol=6 hr = 13 weeks, suggesting filtration but not degradation; amosite: aerosol = 6 hr = 13 weeks, suggesting neither filtration nor degradation. The absence of a filtration effect on amosite fibers might be due to the narrower fiber diameters of that mineral as compared to the MMVFs, which could give long amosite fibers a smaller, hence more respirable, aerodynamic diameter than long MMVF fibers, which might have an aerodynamic diameter large enough to make them less respirable.

Comparison of Two Methods for Measuring Lung Fiber Burden

The literature suggests that destroying the lung to assay retained fibers may change their lengths; Warheit et al. (6) showed that glass fiber dimensions become modified during stimulated tissue digestion, possibly leading to an overestimation of the numbers of fibers/lung (e.g., as a result of
Articles  

Figure 7. Five points on each fiber length cumulative frequency distribution for each fiber type before aerosolization and as retained in lung after 13-week exposure (ashing/SEM, CLSM). (A) MMVF10a; (B) MMVF33; and (C) amosite. Abbreviations: CLSM, confocal laser scanning microscopy; SEM, scanning electron microscopy; MMVF, man-made vitreous fiber. CLSM measurements were in situ three-dimensional measurements on fibers in parenchyma.

Figure 8. Fiber length cumulative frequency distributions of three types of fibers retained in lung after 13-week exposure as determined in situ in three dimensions by confocal laser scanning microscopy. Curve A, man-made vitreous fiber (MMVF)10a; curve B, MMVF33; curve C: long amosite (high dose).

transverse fragmentation). In the present study, the in situ CLSM method provides data on fiber length free from any artifacts due to lung destruction. With MMVF10a, in situ CLSM produced a postretention FLCFD that overlaid that from ashing/SEM analysis (Figure 6A), suggesting that ashing did not fragment retained MMVF10a fibers significantly. Results from MMVF33 run counter to the breakage-during-ashing hypothesis; the FLCFD from in situ CLSM shows a greater proportion of short fibers than that from ashing/SEM. We cannot yet account for this discrepancy. The amosite results, however, were consistent with breakage during ashing; the FLCFD by in situ CLSM showed a greater proportion of long fibers than the FLCFD by ashing/SEM, and the numbers of actual fibers/lung determined by SEM were approximately 250% greater than those determined by in situ CLSM. However, other differences in the two analytical methodologies may offer a better explanation for the discrepancies in amosite counts.

With only two exceptions (MMVF10a and MMVF33 fibers <5 μm long), fiber burdens estimated by ashing/SEM were greater than those estimated by in situ CLSM by factors of 1.4–9.1 (Table 2). The largest difference in counts (ninefold) between the two methods was for amosite fibers <5 μm in length (Table 2). At least two parameters, fiber diameter and fiber distribution throughout the lung, could account for the reduction in the numbers of fibers detected by the in situ CLSM method relative to the ashing/SEM method. The images in this study were collected using the CLSM in the reflected light mode and had pixels approximately 0.51 μm on a side. Fibers with diameters ≤0.5 μm may not have reflected enough light to raise the intensity of the image’s corresponding pixel(s) above noise level. This could explain the especially large (two- to ninefold) discrepancies between SEM and CLSM counts for amosite, which had smaller mean diameters than the two MMVF fibers. Additionally, in situ CLSM would detect fewer fibers than ashing/SEM if fiber deposition was not homogeneous through the lung. For example, during sparse random imaging of a lung sample by CLSM, it would be relatively rare to encounter fibers clumped in a few dense local concentrations; however, with ashing/SEM analysis, all fibers—whether clumped or isolated—would be equally likely to be counted. Finally, fibers approximately 5 μm long can be phagocytized by alveolar macrophages and cleared to pulmonary lymph nodes, where they may remain for an indefinite period (25). Fibers within lymph nodes would have been detected by ashing/SEM, but in the present study no lymph nodes were examined by in situ CLSM. This difference between the two methodologies may adequately account for many of the observed discrepancies.

Evaluation of Analysis of Retained Fibers by in Situ CLSM

There is room for technical improvements, notably in the increase and calibration of detection sensitivity. Effort also must be made to ensure that fibers are not unduly disturbed during tissue preparation and embedding and that tissue configuration remains reasonably unchanged. In addition, in situ analyses are time- and labor-intensive, although advances in speed of image acquisition and subsequent computation continue to ameliorate this situation.

To its credit, in situ CLSM analysis brings several advantages to the study of fiber toxicity models. Because retained fibers may be visualized largely undisturbed, their locations relative to cells and surrounding tissue can be studied and
understood. Some progress toward this understanding can be made using the ashing/SEM technique; lungs may be lavaged, and the lavage fluid may be centrifuged so as to pellet the cells (primarily alveolar macrophages) but not the free extracellular fibers, which then remain in the supernatant. Ashing/SEM may be used to examine the fibers in the lavaged lung and the lavage pellet, producing estimates of the fibers that penetrated lung tissue and those which macrophages had engulfed or attempted to engulf. However, this technique is vulnerable to the uncertain nature of lavaging; it is not clear, for instance, whether all fibers in alveolar air spaces would be swept out into the lavage fluid or whether they could remain trapped.

Additionally, ashing/SEM still does not provide information on where and how fibers and tissue interact or on whether macrophages engulf fibers successfully. In situ CLSM analysis, as shown here, does provide information on these points.

Three-dimensional reconstructions of fibers and tissue can be particularly informative about the modes by which fibers mechanically damage tissue and the mechanisms the organ uses to respond to the insult; in the present study, for example, fibers penetrated septa and airway walls and were partially or entirely engulfed by phagocytes. It would seem possible, also, to use three-dimensional microscopy to study whether fibers and their fragments migrate within tissue or are carried within macrophages, possibly to be cleared from the lung.

Table 2. Fiber fractions and fiber burdens by length class

| Fiber burden (number of fibers/lung) × 10^6 | Fiber fractions | 
|------------------------------------------|-----------------|
| MMVF10a                                  |                |
| Total                                    | 24.8           |
| >0–5                                     | 9.35           |
| >5–10                                    | 9.33           |
| >10–15                                   | 3.4            |
| >20                                      | 1.0            |
| WHO (>5)                                 | 15.4           |

| MMVF33                                  |                |
| Total                                    | 32.8           |
| >0–5                                     | 10.8           |
| >5–10                                    | 10.8           |
| >10–15                                   | 4.73           |
| >20                                      | 2.9            |
| WHO (>5)                                 | 22             |

| Long amosite (high dose)                 |                |
| Total                                    | 68.3           |
| >0–5                                     | 17             |
| >5–10                                    | 23.8           |
| >10–15                                   | 11.9           |
| >20                                      | 9.04           |
| WHO (>5)                                 | 51.3           |

Abbreviations: SEM, scanning electron microscopy; CLSM, confocal laser scanning microscopy; MMVF, man-made vitreous fiber; WHO, World Health Organization. Length distributions of MMVF10a, MMVF33, and long amosite fibers (high-dose regimen) fibers, when aerosolized but before inhalation, and fiber burdens and fiber length distributions at 13 weeks exposure, as obtained by ashing/SEM and in situ CLSM.

*Results from Müller/18.

*Values are ratios of fiber burdens [retained (SEM)/retained (CLSM)].

Table 3. Fiber length densities in hamster lungs

| Fiber types | Nfieldsof-view | Nfiber profiles | Lf | 
|-------------|----------------|----------------|----|
| MMVF10a     | 87             | 359            | 30.5 |
| MMVF33      | 78             | 267            | 25.3 |
| Aamosite (low dose) | 50 | 135 | 20.0 |
| Aamosite (high dose) | 106 | 1,170 | 81.6 |

Abbreviations: Lf length density of fibers in lung; MMVF, man-made vitreous fiber. Length densities of fibers (four dosage regimen) retained in hamster lungs after 13 weeks exposure, as determined by stereological analysis of optical sections obtained by confocal laser scanning microscopy.

*Millimeters per cubic micrometer.

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