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BY AIR MAIL

E.B.A. Co. Ltd. No. 1761

E. Muehleck, Esq.,  
Messrs. Reasbey & Mattison Co.,  
Ambler,  
Pennsylvania,  
U.S.A.

4th June, 1951.

Dear Mr. Muehleck,

Experimental Studies of Asbestosis

The paper by Vorwald, Durkan and Pratt on "Experimental Studies of Asbestosis" has now been studied very carefully by our Factory Medical Officer and others interested in the problem of asbestosis, who report to me that it represents the published version of the earlier Saranac report, which was circulated confidentially. As you know, we had a copy of the latter a little time ago. We find that there is nothing new in substance in the present publication, which constitutes none-the-less an extremely valuable contribution to the study of asbestosis, particularly as its contents can now be revealed to other interested parties.

It is our intention to make copies available to pathologists concerned with post-mortems in presumed asbestosis cases, and for that reason I should be obliged if you would kindly send me another dozen copies.

You ask me for a copy of any written summary or commentary which might be prepared on the present publication, but I am sending instead, as I think it might be more valuable, a copy of the report which was prepared by our Factory Medical Officer about a year ago summarising and comparing work on asbestosis which has been carried out at Saranac and other centres. This should give you a fairly complete picture of the up-to-date position in regard to the study of asbestosis. I should add that we are now sponsoring further work by Dr. Beattie on the question of particle size.

Yours sincerely,

PLAINTIFF'S  
EXHIBIT

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EXHIBIT CC

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30  
COPY LETTER FROM  
TURNER BROTHERS ASBESTOS COMPANY LTD.

Mr. W.H.F. Shepherd.

From

ROCHDALE

RGS/DC

To

E. Muehleck, Esq.,  
Messrs. Keasbey & Mattison Co.,  
Ambler.

T.B.A. No. 1269

Ref.

7th August, 1951.

BY AIR MAIL

Dear Mr. Muehleck,

Experimental Studies of Asbestosis

I have today duly received the 12 copies of the Saranac paper which you mentioned in your letter of 11th June as having been despatched by sea mail. Thank you very much.

Yours sincerely,

*[Handwritten signature]*

201/22M

063 04 00011

File copy

G L  
E. Muehleck, Esq.,  
Keasbey & Mattison Company,  
Ambler,  
Pa.

PER AIR MAIL

JAS/MB T&N.No.4686

16th May 1951.

Dear Mr. Muehleck,

re Asbestosis Report

In Mr. Shepherd's absence, I am writing to acknowledge receipt of your letter of 11th May enclosing copies of your letters dated 10th May to Mr. R. Starkey and Mr. C. H. Jackson, together with a copy of the Saranac Laboratory's report entitled "Experimental Studies of Asbestosis". I note that when you were in Manchester arrangements were made for copies of the report to be made available for T.A.C.Co.Ltd., W.C.Co.Ltd. and T.B.A.Co.Ltd., and we will, of course, let you know if additional copies are required.

Yours faithfully,  
46/

## EXPERIMENTAL STUDIES OF ASBESTOSIS

ARTHUR J. VOIRWALD, Ph.D. (Paris), M.D.

THOMAS M. DURKAN

AND

PHILIP C. PRATT, M.D.

SARANAC LAKE, N. Y.

**ASBESTOSIS** is a form of pneumoconiosis resulting from prolonged inhalation of asbestos dust. The name "asbestos," literally "unburnable," is not that of a specific mineral but is a term applied to a number of different minerals whose characteristic feature is a structure composed of long, parallel, flexible fibers. This structure is unique because the fibers are capable of repeated longitudinal subdivision to units of molecular proportions. In length the fibers vary from a few microns to 6 or more inches (15 or more cm.). Some varieties are stiffer than others, but many are sufficiently flexible to be spun into yarn and woven on modified textile machinery.

The asbestos minerals are silicates of variable composition and belong to the serpentine and the amphibole groups. Listed below are the more common varieties.

Amphibole group: actinolite, anosite, amphibole, anthophyllite, crocidolite and tremolite.

Serpentine group: chrysotile.

The bulk of the asbestos of commerce is chrysotile,  $3\text{MgO} \cdot 2\text{SiO}_2 \cdot 2\text{H}_2\text{O}$ , which is mined on this continent principally in the Thetford region of the Province of Quebec, Canada, and in Vermont. Crocidolite and anosite also are used commercially but in much smaller amounts. Chrysotile occurs as veins in serpentine, a mineral of similar chemical composition, which exists in massive form and is made up of microscopic fibers without the parallel orientation characteristic of chrysotile. The massive, bluish black serpentine, which is smooth and soapy to the touch, is traversed by veins of fibrous chrysotile varying in width from a barely perceptible line to 6 (15 cm.) or more inches. The fibers run across the vein and not lengthwise with the formation.

From the Saranac Laboratory of the Edward L. Trudeau Foundation.

This series of studies of asbestosis, initiated at the Saranac Laboratory more than twenty years ago by the late Dr. LeRoy U. Gardner, director of the Laboratory, was nearly completed at the time of his death in October 1946. Although partial reports and informal reviews of some of the experiments had been given from time to time by Dr. Gardner, this paper presents for the first time a complete survey of the entire experimental investigation.

PK 25-3 (43 pages)

Attention is directed to the mineral brucite,  $Mg(OH)_2$ , which is often found in the same formations with serpentine and clay-solite and may be fibrous in structure. Except for the manufacture of magnesium, brucite has no commercial value at present because its fibers are not sufficiently flexible to be used in textiles, but they are capable of repeated longitudinal subdivision. Unlike other asbestiform minerals, brucite is not a silicate, and for this reason it has been a valuable tool in an experimental evaluation of the action of fibrous minerals on lung tissue.

#### EXPERIMENTAL ASBESTOSIS

For many years studies<sup>1</sup> have been carried on at the Saranac Laboratory in an investigation of the cause, nature and development of asbestosis. The present paper is devoted to experimental asbestosis,

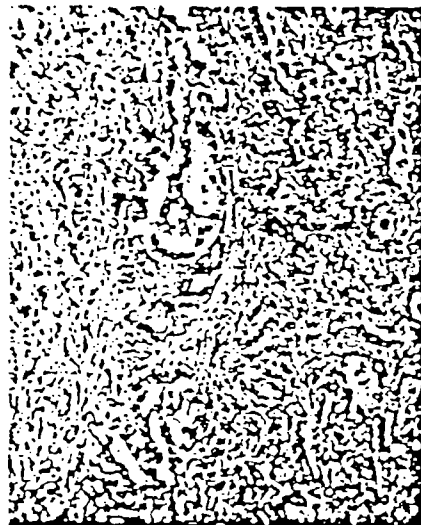


Fig. 1.—Human asbestosis (P-36-134). The photomicrograph reveals a bronchiole (right center) with a smooth muscle bundle at its inferior margin and with an extensive zone of collagen deposition largely obliterating the surrounding alveolar structure. The black body, a wax repository, containing an internal fragment. Asbestos bodies are present but are not apparent at this magnification ( $\times 200$ ).

and in it are described the experiments made on animals with various kinds of asbestos dust. Another report, to be prepared and issued at a future date, will be concerned with human asbestosis and will cover the health aspects of workers who have been exposed to asbestos dust in an industrial environment.

Although in man asbestosis is a chronic disease with diffuse pulmonary fibrosis which requires years to develop, it is possible to reproduce

1. (a) Gardner, E. U., and Cummings, D. E.: Studies on Experimental Pulmonary Infection. VI. Inhalation of Asbestos Dust. Its Effect upon Primary Tuberculosis Infection. *J. Indust. Hyg.* 12: 63 and 97, 1931. (b) Gardner, E. U.: Chrysotile Asbestos as an Indicator of Subtle Differences in Animal Tissues. *Am. Rev. Tuberc.* 42: 562, 1942.

in one or more species of animal characteristic tissue changes which are similar to the lesions of human asbestosis (fig. 1). Since the life span of the experimental animal is relatively short, it is not possible to produce the characteristic lesions in animals under conditions identical with the usual industrial environment. Consequently, to obtain a complete evaluation of the tissue response to inhaled particulate and fibrous material, it is necessary to accelerate the reaction by employing higher concentrations of dust than would ordinarily be encountered in industry. While conditions of exposure are thus different, the information yielded by animal experiments is invaluable in furnishing a better understanding of the reaction of the human organism to inhaled asbestos dust.

#### EXPERIMENTAL METHODS

For investigating the tissue reactions of experimental animals to the various asbestos minerals, two types of technique have been employed, namely, the inhalation method and the injection method. In inhalation experiments, groups of animals—up to 100 or more guinea pigs and sometimes smaller numbers of rabbits, cats, dogs, rats or mice—are kept for eight hours a day in a cubical dust room, 8 ft. (2.5 M.) in dimension, in which a cloud of asbestos dust is maintained by a rotating paddle in a dust hopper.<sup>16</sup> At intervals during the experiment a few animals are killed and the tissues examined to determine the nature and the extent of the dust reaction. Some animals are exposed for periods up to three years. The injection experiments are used to determine in as short a time as possible whether or not a particular dust has a potential capacity to produce inflammatory reaction when in direct contact with tissues of the body. The method involves injecting the dust, either dry or suspended in fluid, into the animal by the intravenous, the intraperitoneal, the intratracheal or another route.

Long term inhalation experiments furnish information on which great reliance is placed when estimating the degree to which a dust might constitute a respiratory hazard to industrial workers. Even though an atmospheric dust may be potentially dangerous, as indicated by injection experiments, only inhalation procedures will reveal whether the dust can be inhaled, pass the natural defense barriers of the body and reach the pulmonary tissue in quantities sufficient to cause damage. Injection methods are useful, however, because they make certain that contact occurs between the dust particles and tissues and because they allow accurate estimation of the dosage and of the potential capacity of that dose to produce reaction. The intratracheal method is particularly valuable when one is dealing with fibrous minerals like asbestos, since it permits observation of the effect of the fibers on pulmonary tissue.

#### TOXIC SIGNIFICANCE

Unlike free silica, asbestos does not produce specific effects in all organs of all species of animals. The comparative data presented in table I are based on completed observations and therefore differ slightly from a preliminary report.<sup>16</sup> Fine quartz introduced into various organs

of various animals: Guinea pig, rabbit, rat, mouse, cat, dog, chicken and even tadpole eventually will produce silicotic nodules but at different rates. Similar introduction of long fiber asbestos has resulted in a fibrous reaction in the lung and, to a lesser extent, in the peritoneum but not in other organs of the guinea pig; the rabbit, the cat and the white rat. In our experience the lungs of the dog and the white mouse failed to respond with fibrosis, although Schuster<sup>2</sup> has reported such changes in a dog that lived in an asbestos-fabricating plant. This variation in species and in organ susceptibility is yet to be accounted for; it is presumed that in the susceptible animals the greater reaction of the lung to asbestos, far exceeding the reaction of other organ tissues, is due principally to the greater vascularity of the lung.

#### PATTERNS OF REACTION OF ASBESTOS

Experience has demonstrated that most of the nonfibrous dust particles inhaled into the lungs of man and animal are 10 microns or less

TABLE 1.—Reaction to Long Fiber Chrysotile in Lungs of Man and Other Species of Animal

| Species     | Route of Exposure        | Fibers* | Asbestos Bodies   |
|-------------|--------------------------|---------|-------------------|
| Man         | Inhalation               | ++      | Abundant          |
| Guinea pig  | Inhalation and injection | ++      | Abundant          |
| Rabbit      | Inhalation and injection | +       | None and atypical |
| Cat         | Inhalation and injection | +       | None and atypical |
| White rat   | Inhalation and injection | +       | Very rare         |
| White mouse | Inhalation               | 0       | None and atypical |
| Dog         | Inhalation               | 0       | None              |

\* The symbols 0 to ++ refer to the degree of reaction.

in maximum dimension. Larger particles apparently do not gain access to the lungs, because, first, large particles settle in air so rapidly that few remain suspended in the atmosphere to be inhaled and, second, large particles are more effectively removed by the protective mechanisms of the upper respiratory tract. In the case of fibrous materials these factors have less influence and fibers 100 and even 200 microns, in length have been found in the terminal air spaces of human lungs. In small laboratory animals exposed to asbestos dust the maximum length of fiber found in the lung rarely exceeds 10 microns.

A large proportion of nonfibrous particulate dust inhaled into the lung is found in the terminal air spaces (alveolar ducts, alveoli, alveoli) in all parts of the organ; in contrast, inhaled asbestos fibers are first discovered in the respiratory bronchioles. These small passages are immediately distal to bronchioles lined by ciliated epithelium.\* Their

\* Schuster, N. H.: Pathology of Asbestos in a Dog. J. Path. & Bact. 34 (p. 275), 1931.

3. Vercauthe, A. J.: Variations in Individual Susceptibility to Industrial Dusts Inhaled into the Lungs. Am. Rev. Tuberc. 62: (111), 1950.

4. Miller, W. S.: The Lung, Springfield, Ill., Charles C. Thomas, Publisher 1937.

own essential lining is a low cuboidal type of epithelium but, as their name implies, they actually function in respiration through lateral alveoli distributed along their walls. Either these alveoli or the abrupt change in the character of the lining epithelium, or the small diameter of the respiratory bronchiole, or the combination of all three factors is responsible for retention of the fiber at this site. Only after asbestosis is well established are appreciable numbers of fibers seen in the more peripheral air spaces. Further explanation is required to clarify this observation.

#### RATE OF TISSUE REACTION TO ASBESTOS FIBERS

The affected tissues react much more rapidly to asbestos than to quartz dust. For example, in rats receiving asbestos fibers by intratracheal injection fibrosis of a characteristic type is visible as early as one month after injection; for quartz dust the latent period is two months or more. Thus, the development of nodular fibrosis due to inhaled silica lags behind the deposition of dust to a greater extent than does the evolution of the diffuse reaction to asbestos. This results in a difference in the degree of progression which follows termination of exposure to dust. For example, on discontinuance of exposure the nodules of silicosis become larger, to a limited extent, for a considerable period of time, whereas the fibrosis of asbestosis increases for only a short time. Subsequently, the asbestotic fibrous tissue contracts; this process often distorts the adjacent pulmonary tissue and may, as a result, progressively interfere with cardiorespiratory function.

#### ASBESTOS BODIES

The peculiar structure known as the asbestosis body or "curious body" is a specific concomitant of asbestosis.<sup>5</sup> The typical body is a golden yellow, beaded or beaded rod, which may be either straight or curved (fig. 2). Often one or both ends are bulbous like a dumbbell. The bodies vary considerably in length, and dimensions up to 250 microns have been recorded.

It is believed that asbestosis bodies are inhaled fibers on which protein and iron pigment of tissue origin have been deposited.<sup>6</sup> Glyne observed reproduction of these bodies in guinea pigs nine months after subcutaneous injection of fibers rendered free of iron. The bodies are abundant in man and in the guinea pig (table 1) but are much larger in the former, probably because the larger-sized air passages admit fibers of greater dimension. In guinea pigs they form after about 70 days

5. Glyne, S. R.: (a) The Formation of the Asbestosis Body in the Lung. Tubercule 32:398, 1931; (b) The Asbestosis Body. Lancet 1:1351, 1932. (c) Gardner and Cummings.<sup>10</sup>

6. Lynch, K. M., and Smith, W. A.: Asbestosis Bodies in Sputum and Lung. J. A. M. A. 95:659 (Aug. 30) 1930. Simon, F. W., and Strachan, A. S.: Asbestosis Bodies in the Sputum: A Study of Specimens from 50 Workers in an Asbestos Mill. J. Path. & Bact. 34:1, 1933. Gardner and Cummings.<sup>10</sup> Glyne.<sup>5</sup>

of contact with the tissue. In fact, talc and mica are found in the lungs. In show an atypical coating after much longer residence in the lungs. In rats the bodies are rarely seen, and in dogs none could be found. Although the evidence is incomplete, it appears that the formation of the asbestos body prevents the fiber from damaging the tissue. Many of the points mentioned above will be elaborated on in subsequent para-

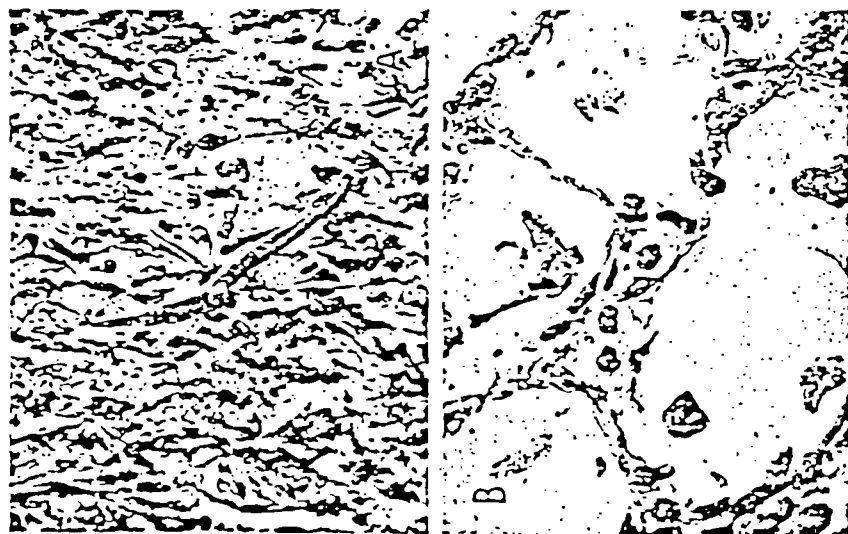


Fig. 2—A, human asbestos body. This collection of asbestos bodies was found in the lung shown in figure 1. The usual variations in size and configuration are represented ( $\times 400$ ).  
B, guinea pig asbestos body. This one is similar to some of those shown in A ( $\times 400$ ).

graphs dealing with the actual experiments. For presentation our investigation is divided into two sections, one dealing with inhalation experiments and the other with injection experiments.

## INITIAL INHALATION EXPERIMENTS

Four large scale inhalation experiments have been conducted in this laboratory with various forms of asbestos dust. In each of these investigations, more than 100 animals were used, and the experiments were carried on for periods ranging from two to more than five years. The four kinds of asbestos dust employed are designated as King's floats, short fiber, 100 per cent ball-milled, and long fiber asbestos dust.

### KING'S FLOATS ASBESTOS DUST

The first inhalation experiment conducted at the Surinac Laboratory with asbestos dust was begun in 1928. Animals inhaled the dust for

TABLE 2.—Chemical Analysis of Asbestos Dusting Materials

| Type of Asbestos | SiO <sub>2</sub> | Fe <sub>2</sub> O <sub>3</sub> | Al <sub>2</sub> O <sub>3</sub> | Cr <sub>2</sub> O <sub>3</sub> | H <sub>2</sub> O | CaO   | MgO   | Na <sub>2</sub> O | K <sub>2</sub> O | CO <sub>2</sub> | Total  |
|------------------|------------------|--------------------------------|--------------------------------|--------------------------------|------------------|-------|-------|-------------------|------------------|-----------------|--------|
| King's floats    | 34.25            | 8.94                           | •                              | •                              | 0.47             | 11.24 | •     | •                 | •                | 12.71           | 97.33  |
| Short fiber      | 31.15            | 9.07                           | 1.40                           | 0.15                           | 0.08             | 0.45  | 15.46 | 0.14              | 0.70             | 0.88            | 100.11 |
| Long fiber       | 34.10            | 8.11                           | 0.16                           | •                              | 0.05             | 0.31  | 10.14 | 0.04              | 0.23             | 1.00            | 99.16  |

• Not determined.

TABLE 3.—Petrographic Analysis of Asbestos Dusting Materials

King's floats. The approximate composition, based on particles (except chrysotile smaller than 10 microns) as determined by percentage obtained from particle counter, was chrysotile 10, actinolite 10, amphibole 10, quartz 10, talc 10, other minerals 10. For chrysotile, there up to 100 microns long were included.

Short fiber 1. The material, before being ball milled, contained a preponderance of fibrous chrysotile and plagioclase amphibole. The approximate composition, by percentage, was actinolite 11, actinolite 22, mesophyllite 10, quartz 2, brucite 4, other minerals, including dolomite, actinolite and tremolite, 11.

Long fiber 1. The material consisted principally of the fibrous asbestos mineral chrysotile. Short of non-persistent fibers 1 to 15 microns in diameter and up to 50 microns in length were present. The approximate composition, by percentage, was chrysotile 73, actinolite 11, mesophyllite 4, brucite 2, other minerals, among which were calcite and chlorite and micaceous minerals, 2. Only a trace of quartz was observed.

The analysis of the King's floats asbestos, made by Dr. C. S. Hurlbut Jr., of Harvard University, has been reported elsewhere (Hurlbut, C. S., Jr., and Williams, G. B.: The Mineralogy of Asbestos, Part 1, *United States Geological Survey Bulletin* 1030, 1930). The analysis of the short fiber asbestos and the long fiber asbestos the petrographic analysis was supplemented with a very diffraction resolution.

periods up to 33 months. Some guinea pigs with six and nine months' exposure lived for an additional three years after cessation of their exposure. A preliminary report<sup>10</sup> presented observations after 29 months of exposure. At that time observations covered a period of only 2 1/4 years and the conclusions as to the ultimate effects of inhaled asbestos dust were provisional. Those conclusions are substantiated by results of the completed study, which is reported as follows.

Composition and Atmospheric Concentration of the Dust.—The dusting material, a commercial variety of asbestos known as King's floats, was composed of short fibers, ranging in length from 1 mm. to 1 micron or less, and of particles which also varied in size. It was obtained from the Thetford, Quebec, plant of the Asbestos Corporation of America, and analyses (tables 2 and 3) revealed that the amount of fibrous chrysotile was only 14 per cent, a rather low value.

were phagocytosed, and many of them were carried into the wall by migratory leukocytes. Mononuclear leukocytes attracted to the area caused an appreciable thickening of the bronchial wall. After 16 months a delicate fibrosis made its appearance. The process evolved gradually, and the number of fine intercellular collagenous fibers steadily increased. As this fibrous deposit contracted, it partially closed and

TABLE 4.—Summary of Substation Inspection, Abatement, and

[illegible]

\* The children's eyes were deflected with low magnitude by stimuli of horizontal location. This confirms the results of several previous studies, but they

Results of the investigation, briefly summarized in table 1, show that inclusion of King's blood vessels did produce a typical pericardial-in blood in Queen's but not in rabbits or rats.

Reaction in Normal Guinea Pigs.— Guinea pigs including this, died for periods up to 33 months after a characteristic fibrosis occurring in connective tissues about the respiratory bronchioles. During this exposure the peripheral alveoli were not involved. The particulate elements of the dust were transported through the lymphatic system to the bronchial nodes, causing no significant reaction in either the lymph nodes or the bronchial nodes. The fibrous elements retained in the alveoli were seldom detected in the bronchial nodes.

After exposure of approximately a year a small amount of cellular reaction had been produced about many respiratory bronchioles (fig. 3A). As more dust was inhaled, it continued to accumulate in the same location, and later stages of the disease (fig. 3B) consisted of extensions of the original lesions.

Apparently, the inhaled fibers were caught in the pocket like alveoli that are covered from the lateral walls of the respiratory bronchioles. There they

7. Fisher, W. B.; Houtz, R. L.; Dudley, A., and Matthews, J. E. *Abstracts of The Collection and Counting of Antibody Data Presented in Abstracts Published in the Proceedings of the 1994 Annual Meeting of the American Society for Cell Biology*, 1994.



Fig. 3—King's sham isolation experiment: *A*, lung of a guinea pig with 12 transplanted tubercles; it includes a respiratory bronchiole, at the left, branching and dividing into alveoli; *B*, lung of a guinea pig with 12 transplanted tubercles; it includes a respiratory bronchiole, at the left, branching and dividing into alveoli; *C*, lung of a guinea pig with 26 months' exposure. The field includes a bronchiole, at the center, with alveoli on either side; the alveoli are filled with tubercles; *D*, lung of a guinea pig with 26 months' exposure, showing the walls of alveoli at right angles to each other, with tubercles lying along these alveoli. This is the so-called "atuberculation" appearance (see text).

distorted the alveoli, and with this change the alveoli became lined with cuboidal cells. The result was an adenoma like appearance which frequently accompanies

characteristic hyaline reaction resulting from many causes. Wilby<sup>9</sup> described a similar structure in the lungs of guinea pigs inhaling asbestos vermiculite. The longer asbestos exposures resulted only in more thickening of the walls of the air spaces, largely due to an increase in the amount of fibrosis. The fibrous tissue always remained cellular and failed to show the hyalinization characteristic of silicosis.

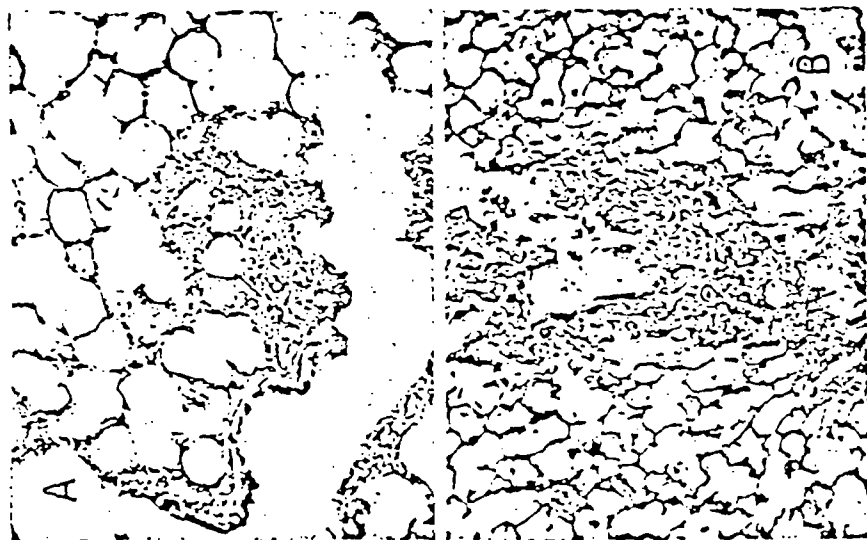


Fig. 4—King's fibrosis inhalation experiment. A, lung of a guinea pig with six months' dust exposure followed by 35 months' inhalation of normal air. The reaction is rather slight, but distinct fibrosis is present (X 200). Note that 28 months of continuous exposure (fig. 3B) produces much more extensive reaction. B, lung of a guinea pig exposed to the asbestos dust for nine months and having fibrotic reaction in nodules at 37 months. The reaction shown is more than that in A but much less than the reaction in figure 3B (X 200).

<sup>9</sup> Wilby, H. V., and Rudasew, P. Tumor-like Structures in the Lungs of Guinea Pigs Artificially Exposed to Fibrous Dust, *Am. Rev. Tuberc.* 47:268, 1946.

Asbestos bodies (fig. 2B), first seen in the lungs of the guinea pigs that had inhaled dust for about two months, became more numerous and more distinctly segmented with increasing exposure.

The reaction produced in guinea pigs exposed for six and nine months did not progress significantly during a subsequent period of 35 and 37 months when the animals lived in a normal atmosphere (fig. 4). Between eight and 11 months after exposure ceased, the cellular reaction in the lung had been completely replaced by thin strands of fibrous tissue. At later periods the scar tissue was less in amount, but in the last animal killed, 37 months after discontinuing dust exposure, some fibrosis was still visible.

**Reaction in Guinea Pigs Infected with *Tubercle Bacilli* at the Onset of Dust Inhalation.**—Of the group of 40 guinea pigs infected with attenuated tubercle bacilli, R<sub>1</sub> strain,<sup>8</sup> at the time that dust exposure was begun, 31 died or were killed before the completion of two years of the exposure and were reported in the paper by Gardner and Cummings.<sup>8</sup> Seventeen of these died from intercurrent pneumonia. Briefly, the results were as follows: Ten revealed some evidence of spread of the tuberculous process (fig. 5A); in 6 of these it was confined to the lungs, and in the other 4 the abdominal viscera also were involved. Extension of the infection was first seen after seven months of dust inhalation; during the next 20 months more than half of the animals showed actively spreading tuberculosis, and in 3 of them small cavities had developed. During the last eight months no animals exhibited any evidence of active infection although in half of them the healed fibrous scars of previous spreads were obvious. The scars were more extensive than is characteristic of either tuberculosis or asbestosis alone.

The nine animals which were still alive after two years of dust exposure were killed at intervals during the following year. In four of them the primary foci of infection were healed with fibrosis and even calcification, and there was no evidence of progression (fig. 5B). In the remaining five the tuberculous foci showed evidence of having previously spread locally; in four of them, by the time of autopsy, the foci were healed with excessive fibrosis; in the fifth animal there was a generalized chronic tuberculous pneumonia in one lobe, and in the other lobes there were isolated primary tubercles, which were still active but had not spread.

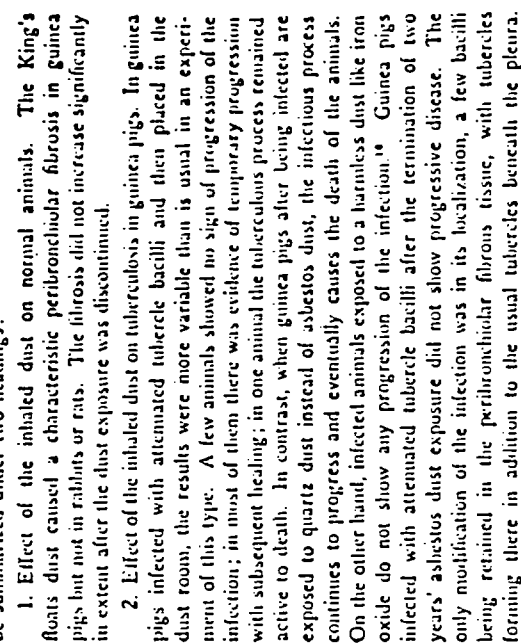
**Reaction in Guinea Pigs Infected with *Tubercle Bacilli* After Establishment of Asbestosis.**—Twelve guinea pigs, after inhaling King's fibrous asbestos dust for 26 months, were infected with tubercle bacilli and then removed to normal air. Six of these animals died within seven weeks, five from intercurrent non-tuberculous infection. The remaining six animals were killed at intervals up to 14 months after infection. The subpleural tubercles were no more numerous in the dusted animals than in the non-dusted controls, but a considerable number were found in the depths of the lung about foci of asbestosis. The tuberculous component of the combined reaction showed only slight local extension about lesions in the lungs and tracheobronchial lymph nodes. Cavitation was found in tubercles 1½ months old, but by 3½ months it had completely disappeared, leaving only scar tissue. Foci of fibrosis still persisted in the last animal, which was killed 14 months after infection.

**Reaction in Rabbits.**—Rabbits exposed to the asbestos dust for periods up to 19 months showed a foreign body type of reaction of low grade, but no fibrosis. Although their lungs contained particulate elements of the dust, fibers were not present, indicating that the upper respiratory mechanism of the rabbit is adequate to exclude fibrous foreign bodies. Two rabbits, after inhaling dust for six and 19

<sup>8</sup> Strecker, W. Jr., and Gardner, L. U.: R. Strain of *Tubercle Bacillus*: Its Dissemination and Virulence of Variants in Normal and Silicotic Guinea Pigs, *Am. Rev. Tuberc.* 44:51, 1946.

their lungs. In a few of the rats, an occasional asbestosis body was discovered, but there was no fibrosis. This phase of the experiment was considered unsuccessful.

**Summary and Interpretation of Inhalation Experiment with King's Floats Dust.**—The findings in the experiment with King's floats dust can be summarized under two headings:



*Comparison and Atmospheric Concentration of the Dust.*—The dusting material for this experiment was the remains of fibers collected in dust bins of an abestos-laminating plant after a carding operation and screened to pass 200 mesh. Since

10 Voraiah, A. J.; Pruthi, P. C.; Durkan, T. M.; Delahant, A. W., and Bailey, D. A. Screening A Plaque Forming Assay Due to the Inhibition of Iron Dost, *Indian Med. J. Surg.* 19 (3) 1959

Shoot Line Asbestos Dust

Since hazardous dusts like quartz are most effective in producing fibrosis when the particles are 3 microns and less in size, an inhalation experiment was performed to determine whether this condition is true for asbestos dust. It was thought that a short fiber asbestos dust consisting almost entirely of fibers and particles smaller than 3 microns would initiate an accelerated tissue response and produce an advanced reaction in a shorter time than did the King's floater dust, which contained fibers from 1 mm to 1 micron and less in length as well as much particulate matter.

*Comparison and Atmospheric Concentration of the Dust.*—The dusting material for this experiment was the remains of fibers collected in dust bins of an abestos-laminating plant after a carding operation and screened to pass 200 mesh. Since

10 Voraiah, A. J.; Pruthi, P. C.; Durkan, T. M.; Delahant, A. W., and Bailey, D. A. Solenophy A Plaque Pneumonia Due to the Inhalation of Iron Dust, *British Medical Journal*, 1970, 1: 539.

Only after exposures had continued for approximately one year was there an appreciable tendency for dust-containing phagocytes to gather into clumps. By 16 months phagocytes had collected about the walls of a few of the respiratory bronchioles which revealed a little proliferation or infiltration of mononuclear cells. There were also some multinucleated cells, but they were of the inert, foreign body type. At 20 to 24 months the cellular clumps were sometimes quite prominent, and sometimes changes in the epithelium resulted in the adenoma-like or "adenomatoid" appearance (fig. 3*B*) previously described in the section reviewing the experiment with the King's flints dust. In most of the subsequent members of the series the reaction remained cellular, but a few exhibited pronounced development of fibrous tissue. In these few members of the series the color of the lung was pale in color and firmness, with no appearance of being hyalinized. Unlike chronic pleurisy was present in a few animals without evidence of pul-

collagen was pale in color and tenacious, with no appearance of being hyalinized. Dense chronic pleurisy was present in a few animals without evidence of pulmonary infarction.

TABLE 6.—*Analyses of Lungs of Guinea Pigs After Prolonged Inhalation of Short-fiber Asbestos Dust*

\* The symbols averaging the group reaction in each group of sub-panels represent merely the relative degree of reaction, ranging from 0 (quadrants) to 4 (the maximum for this experiment). The relationships apply only within this table and cannot be compared with quadrants in other tables.

In the group returned to normal air after 20 months' inhalation of dust, progression of disease was not definitely demonstrated, but neither could it be absolutely disproved, owing to the variability of the response in different animals. The treatments, from mild to severe, occurred sporadically and bore no relationship to the length of time after cessation of exposure. The differences were attributed to variation in individual susceptibility. This view received support from the biochemical analyses (table 6), which revealed comparable amounts of ash and silica in the lungs with widely different amounts of tissue change. For example, the ash

The dust concentration varied during the experiment, the light field counts for atmospheric samples collected inside the animal cages with the impinger apparatus ranging from 81 million to 185 million. The average of counts was 130 million for the first year of the experiment, 134 million for the second year and 140 million for the third year.

TABLE 5.—Summary of incubation experiments with Sheet Pile, *Asiatrya* Pond

• After 24 months the animals were exposed to one per cent lipid with the glucose solution.

• The reaction was generally slow to look blood in the myocardial treated which was infuse with the glucose solution short

• glucose was estimated short there

Four species of animals—gamma mice, white rats, cats and rabbits—were used in this experiment. The results of the dust exposure, summarized in Table 3, are presented in greater detail below.

The type of tissue reaction provoked by the inhaled dusts that substantially differed from that provoked by the inhaled dusts that did not, was essentially the same as that already observed in the experiments with King's dust, as described above. The rate of reaction also was approximately the same, but the extent of involvement was very much less. After 10 to 21 months of exposure only a very small part of the trachea, which generally required microscopic examination for detection, had been involved in the allergic process.

The formation of alysiostichus bodies was at first extremely limited in both groups. After five months' exposure only a very rare short body could be found, usually a single cell. Some of the forest intragular patches were surrounded by yellow alysiostichus having the same color as the alysiostichus body. One year's exposure had permitted an accumulation of many longer filices, a number of which were coated and covered as typical alysiostichus bodies. Most of these were still short enough to be partially or entirely within phragmocytes. By the twentieth month and thereafter they

TABLE 7 ... ANALYSIS of Income of White Kids That Had Included Short-Term Absences Paid

| Location<br>of Mine | Amount<br>of Ore<br>Treated,<br>Long Tons | Total<br>Residue,<br>Long Tons | Duration<br>of Run,<br>Mins. | Amount<br>of Ore<br>Treated,<br>Long Tons | Total<br>Residue,<br>Long Tons |
|---------------------|-------------------------------------------|--------------------------------|------------------------------|-------------------------------------------|--------------------------------|
| A                   | 22                                        | 0.0                            | 0                            | 22                                        | 0.0                            |
|                     | 22                                        | 0.0                            |                              | 22                                        | 0.0                            |
|                     | 22                                        | 0.0                            |                              | 22                                        | 0.0                            |
| B                   | 20                                        | 0.0                            | 0                            | 20                                        | 0.0                            |
|                     | 20                                        | 0.0                            |                              | 20                                        | 0.0                            |
|                     | 20                                        | 0.0                            |                              | 20                                        | 0.0                            |
| C                   | 22                                        | 0.0                            | 0                            | 22                                        | 0.0                            |
|                     | 22                                        | 0.0                            |                              | 22                                        | 0.0                            |
|                     | 22                                        | 0.0                            |                              | 22                                        | 0.0                            |
| D                   | 22                                        | 0.0                            | 0                            | 22                                        | 0.0                            |
|                     | 22                                        | 0.0                            |                              | 22                                        | 0.0                            |
|                     | 22                                        | 0.0                            |                              | 22                                        | 0.0                            |
| E                   | 22                                        | 0.0                            | 0                            | 22                                        | 0.0                            |
|                     | 22                                        | 0.0                            |                              | 22                                        | 0.0                            |
|                     | 22                                        | 0.0                            |                              | 22                                        | 0.0                            |
| F                   | 22                                        | 0.0                            | 0                            | 22                                        | 0.0                            |
|                     | 22                                        | 0.0                            |                              | 22                                        | 0.0                            |
|                     | 22                                        | 0.0                            |                              | 22                                        | 0.0                            |

• Supply of credit to the food industry

TABLE 8—Average Values of 4th and Total Values for Twenty of White Kents Including Current Basis for Current Periods (Twenty Only, if about Included Twenty Only)

| Date | Amount of Ashes Used in Fuel |       |       |       | Total Ashes, % of Total Fuel |       |       |       | Total Sulfur, % of Ash |       |       |       |
|------|------------------------------|-------|-------|-------|------------------------------|-------|-------|-------|------------------------|-------|-------|-------|
|      | W.B.A.                       |       | Fuel  |       | W.B.A.                       |       | Fuel  |       | W.B.A.                 |       | Fuel  |       |
|      | Grass                        | Grass | Grass | Grass | Grass                        | Grass | Grass | Grass | Grass                  | Grass | Grass | Grass |
| 1    | 2.3                          | 2.9   | 2.9   | 0.92  | 0.31                         | 0.75  | 0.46  | 2.6   | 11.7                   | 3.9   | 7.6   | 3.9   |
| 2    | 4.8                          | 5.6   | 5.6   | 0.92  | 0.4                          | 0.12  | 0.07  | 2.5   | 11.6                   | 3.6   | 7.9   | 4.0   |
| 3    | 3.5                          | 3.1   | 3.9   | 3.4   | 0.04                         | 3.6   | 0.11  | 1.6   | 4.5                    | 4.6   | 4.6   | 4.6   |
| 4    | 3.8                          | 4.6   | 4.9   | 3.6   | 0.1                          | 1.6   | 1.00  | 0.22  | 2.9                    | 5.1   | 5.0   | 5.1   |
| 10   | 4.9                          | 7.6   | 14.1  | 0.1   | 0.15                         | 4.6   | 0.19  | 0.75  | 2.2                    | 5.0   | 4.5   | 4.5   |

were relatively numerous although still rare in comparison with the findings in the King's theory experiment.

**Reaction to White Rot.**—Seventy-three white rats were exposed to atmospheric dust for a period of 12 months. During the first 10 months the rats were killed bi-monthly and for the remainder of the experiment at less frequent intervals. Up to eight months the dust cells were solely observed and counted in the smears. Reaction was limited to occasional slight thickening of the septa, no great small accumulations of dust cells. At 10 months there was a migration of early blastema in a few rats, but the change was so slight that it would probably have been overlooked without the change of dust cells which attracted attention to the area. Only 10 animals were exposed for from 12 to 32 months. In each of them the lungs contained minute foci of well-defined foci of granuloma, like that of alveolitis but without alveolar lesion. The lesions, although little else, of a granuloma of 100 diameter or more, consisted of not only alve-

alveolar ducts in which the walls of the associated air spaces were very thick, owing to a swollen collagen framework. Connective tissue and blood-rich, heavily silver preparations revealed complete loss of capillary bed locally. Outside the collagen was a thin layer of epithelial cells. This did not resemble the "adenomatous" change characteristic of guinea pig asbestosis. Near the lesions the air spaces were filled with phagocytes containing gray to yellow particulate dust and a rare, long, uninked asbestos fiber. Careful search failed to reveal even a suggestion of an asbestosis body. Pleurisy was absent. The tracheobronchial nodes showed compact focal collections of mononuclear cells at 12 months and, at 20 months, some diffuse thickening of the reticulum. In a few rats there was definite fibrosis along the margins of the nodes, extending into the mediastinal areolar tissue.

Results of chemical analyses made on the white rats are given in table 7, and the average values have been recorded in table 8 for comparison with similar values for rats inhaling other dusts. It will be noted that the values for asbestos are lower than those for quartz or chert but approximate those for the gypsum-quartz mixture, in which atmospheric agglomeration tended to reduce the amount of dust inhaled. This condition prevailed even though the atmospheric concentration of asbestos dust was essentially the same as that of the quartz, was one-half that of the gypsum-quartz mixture and was one-fifth that of the ferruginous chert. Since the values for asbestos are low, it might be inferred that the total quantity of dust actually inhaled was small or that it had been eliminated from or dissolved within the lungs. Evaluation of these possibilities is not feasible on the basis of the observations derived from this study.

**Reaction in Cats.**—Twenty cats were used in this inhalation experiment with the short fiber asbestos. Eighteen were kept in the dust room continuously until they died of asbestosis, the exposure period ranging from one month to nearly 54 months. The other two were removed to normal air after a short exposure of 31 months; none of these was killed five months, and the other 24 months, later. In general, the tissue response was confined to microscopic foci of fibrosis, which were in the subpleural areas of groups of subpleural alveoli rather than in the peribronchovascular areas. In some animals the change was extensive enough to be visualized on gross inspection of the section. Only in the animal with the longest exposure—54 months—did the roentgenogram reveal definitely abnormal shadows. A roentgenogram made after 30 months revealed no abnormality; after 45 months, a faint mottling could be detected throughout both lungs. At autopsy, nine months later, there was only microscopic fibrosis in the subpleural zone plus heavy lymphocytic infiltration about small bronchioles. Asbestosis bodies were rare. On prolonged search a few yellow atypical bodies, amorphous and without inclusions, were found in two animals exposed for more than a year.

**Reaction in Rabbits.**—Eight rabbits were exposed to dust for periods extending from one to more than six years; the last animal was removed from the dust chamber and left in normal air six months before being killed. There was never enough pulmonary fibrosis to be detected grossly, and there was no chronic adhesive pleurisy. Microscopic evidence of alveolar wall thickening was first detected in one animal after about three years of exposure and was seen in all five animals examined thereafter, including the one removed to normal air. One animal that died of paralysis after only four years of exposure exhibited a reaction visible on gross inspection of tissue sections. The possibility of pulmonary infection in this animal could not be excluded. In another animal dying two years later the alveolar fibrosis was not nearly as obvious or as advanced. Areas of involvement, which were largely visualized because of phagocytic reaction within the air spaces, extended microscopically to become more fibrous with the passage of time, but there was never much entanglement on the lumen of air spaces and the structure of the lung was preserved. Asbestosis bodies were not detected in rabbits that died early in the experiment but were seen in all animals that had been exposed to the dust for more than three years.

**Summary and Interpretation of Inhalation Experiment with Short Fiber Asbestos Dust.** The original purpose of the experiment was to evaluate the role of short asbestos fibers in the genesis of asbestosis. It was felt also that if the tissue reaction to short fibers was not as extensive as believed, that the action of asbestos is in part, at least, a chemical one as postulated for quartz. This experiment, in which the tissue reaction was slower and less extensive than that in the previous experiment with King's flake dust, indicates that the capacity of inhaled asbestos fibers to produce fibrosis is determined primarily by factors not chemical in nature.

On the four species exposed in this experiment, only the guinea pig and to a lesser extent the white rat responded with characteristic peribronchovascular fibrosis. The rat reacted with atypical subpleural fibrosis and the rabbit with only slight peribronchovascular fibrosis.

#### Box-Martin Asbestos Dust

In the inhalation experiment with short fiber asbestos dust a small quantity of unground short fiber asbestos was mixed with the ball-milled product in order to generate a suitable dust cloud. When that experiment failed to produce an accelerated tissue reaction, in conjunction with the response initiated by King's flake dust, it became apparent that the biologic activity of asbestos is not increased by a reduction of fiber size. Thus the possibility arose that the tissue reaction observed was due solely to the relatively few long fibers of the unground asbestos and that the short fibers of asbestos had no more than a very insignificant role in the production of asbestosis, a concept not in accord with previous experiments concerning pneumoconiosis. Consequently another inhalation experiment was started in which only ball-milled asbestos was used.

**Composition and Homogeneity of the Dust.** The dusting material was the ball-milled short fiber asbestos used in the previous inhalation experiment, but unground material was not mixed with it. Owing to the tendency of the material to form small spherules which prevented much of the fibrous portion from floating out of the dusting machine, the dispersal of the dust was not entirely satisfactory. Therefore, after an initial warm-up period of operation, steel wire brushes were attached to the inside surface of the hopper and to the rotating paddle to disintegrate the spherules and release the fibers. This arrangement gave satisfactory results and was used for the remaining 21 months of the experiment.

The composition of the raw asbestos used is shown in Tables 2 and 3. Petrographic and x-ray diffraction examination of atmospheric dust, collected in the dust room with an electrostatic precipitator after the installation of wire brushes, indicated that about 15 per cent of the air-suspended material was chrysotile, and about 85 per cent, serpentine; of the balance, magnetite comprised 10 per cent, hematite 3 per cent, quartz 2 per cent and other minerals 10 per cent. During the seven-month period before the wire brushes were used, the chrysotile content of the atmospheric dust was somewhat lower than 15 per cent, but reliable values were not obtained.

The dust concentration during the first seven months of the experiment was about 100 million particles per cubic foot of air. After the wire brushes were

installed, the dust counts were higher, and the over-all average for the remaining 21 months was about 150 million.

Site-frequency studies of atmospheric dust collected inside the animal cages revealed that nearly 99 per cent of the components suspended in the air could be classified as fibrous; only about 1 to 1.5 per cent was fibers. One third to one half of the fibers were longer than 10 microns, indicating a concentration of long fibers of about 0.8 million. This figure is about one-half the estimated value of 1.4 million for the short fiber experiment.

Guinea pigs, rats and mice were used in the inhalation experiment with the 100 per cent ball-milled asbestos dust. The results are summarized in Table 9.

**Reaction in Guinea Pigs.**—The experiment was started with 100 guinea pigs. As the dust exposure proceeded, there were 39 accidental deaths, 32 of these being due to pneumonia in an epidemic. The 61 pigs remaining exposed to the dust were killed at intervals during exposure, except for 16 guinea pigs transferred to normal air after 28 months of dusting. For the first year of exposure practically the only reaction to the dust was the presence of scattered phagocytes and an occasional minute asbestos body. At 16 and 20 months no gross response was visible on the tissue section, but microscopically peribronchovascular loci of inflammatory cells

Table 9.—Summary of Inhalation Experiment with 100 per Cent Ball-Milled Asbestos Dust

| Nature of Exposure                                      | Animals | Maximum |       | Results                           |
|---------------------------------------------------------|---------|---------|-------|-----------------------------------|
|                                                         |         | Mean    | After |                                   |
| Dust exposure on: 24 guinea pigs                        | 24      | Mean    | Mean  | No appreciable pulmonary reaction |
| out life                                                | 24      | Mean    | Mean  | No suggestion of asbestosis       |
| Dust exposure in: 16 guinea pigs                        | 16      | Mean    | Mean  | No suggestion of asbestosis       |
| level of peribronchovascular loci of inflammatory cells | 16      | Mean    | Mean  | No suggestion of asbestosis       |
| in normal air                                           | 16      | Mean    | Mean  | No suggestion of asbestosis       |

could be seen. At 24 months (Fig. 6A) there was still no change large enough to be seen with a hand lens, although microscopic examination revealed cellular accumulations about terminal bronchioles and many more asbestos bodies, chiefly within cells. The lungs of animals exposed for the full dusting period of 28 months and afterward living in normal air for two months revealed the changes described above and also very slight peribronchovascular fibrosis. For exposed animals living eight months in normal air the findings were similar, but at 12 months three of four animals showed grossly visible characteristic peribronchovascular fibrosis with adenomatoid change (Fig. 6B).

The tracheobronchovascular nodes were essentially normal until exposure had been continued for more than a year and a half. Animals killed at 12 months and at 16 months revealed a few minute collections of phagocytes containing particles but practically no fibers large enough to be recognized as such. After 20 months of exposure many monocytes filled with yellow granules were present. At 30 months there had been a slight increase in reticulum but no fibrosis. No further changes occurred in the nodes. Asbestos bodies were not seen in the nodes of any of the guinea pigs.

Minute asbestos bodies were observed in the lungs as early as three months after exposure began, but they did not become numerous until 16 months had elapsed. The bodies were short and practically all were intracellular, although at 20 months some were long enough to project beyond the cell borders. It is

important to note that in the later months of exposure there was a distinct increase in the number of lung fibers, up to 20 microns in length, in the lungs with the formation of characteristic long asbestos bodies.

Chemical analyses (table III) of the lungs revealed that considerable dust had been retained in the lungs. After 24 months of continuous exposure the average

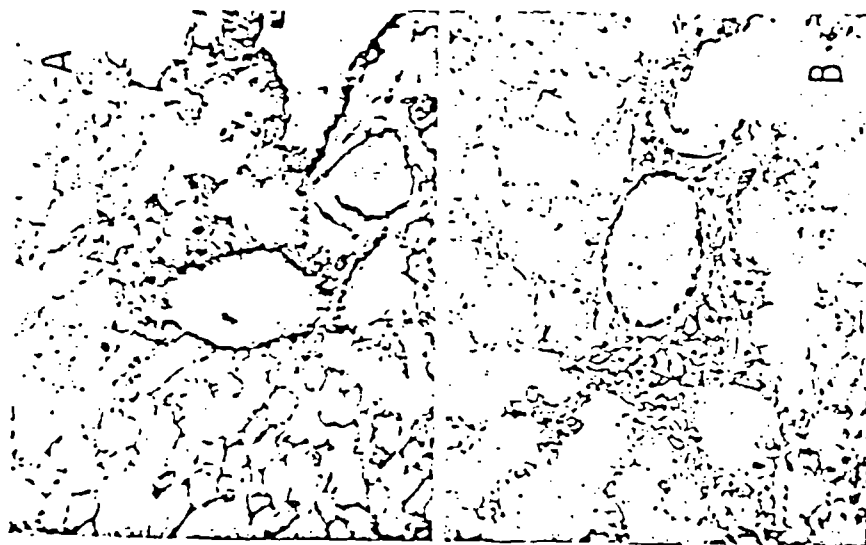


Fig. 6: Ball-milled asbestos inhalation experiments. A: Lung of a guinea pig with 24 months' dust exposure. A bronchiole is shown at the center, with a thick accumulation of pleural cells but without the formation of collagen (C. 200 $\times$ ). B: Lung of a guinea pig with 24 months' dust exposure and then 12 months' inhalation of washed air. The reaction is much like that shown in A, but there is a slight deposition of collagen, most apparent at the left (C. 200 $\times$ ).

value for total silica, per cent of ash, was 25.47. This should be contrasted with the average value of 14.31 (table 6) for animals exposed 24 months to the short fiber asbestos dust.

In view of the high values for silica obtained with the animals exposed to 100 per cent ball-milled dust, it is important to note that their pulmonary response was much less than that of animals exposed for 24 months to the short fiber asbestos in the previous experiment. This again indicates that the biologic activity of asbestos inhaled into the lung is not increased by a reduction in size of the fibers.

**Reaction in White Rats and Mice.**—In this experiment 40 rats were exposed for periods up to 20 months and 24 mice for periods up to 12 months. In neither species did even a suggestion of asbestosis develop, and reaction was limited to phagocytosis of inhaled particles by widely scattered dust cells which remained free in air spaces or were transported to the tracheobronchial lymph nodes. No asbestos bodies were found in the rats, but in the mice there were a very few small, nonhydrated forms within phagocytes.

Table 10.—Analyses of Lungs of Guinea Pigs Exposed to Dust in Inhalation Experiment with 100 per Cent Ball-Milled Asbestos Dust

| Exposure to Dust, Mo.                                       | Period in Normal Air, Mo. | Amt. of Ash, % of Dry Lung | Total Silica, % of Ash | Total Silica, % of Ash | Tissue Reaction* |
|-------------------------------------------------------------|---------------------------|----------------------------|------------------------|------------------------|------------------|
| 1                                                           | 0                         | 1.55                       | 0.21                   | 4.28                   |                  |
|                                                             |                           | 1.30                       | 0.20                   | 7.60                   |                  |
|                                                             |                           | 4.35                       | 0.24                   | 6.60                   | 0                |
| 1                                                           | 0                         | 4.68                       | 0.22                   | 4.90                   |                  |
|                                                             |                           | 1.60                       | 0.14                   | 1.90                   |                  |
|                                                             |                           | 8.05                       | 0.41                   | 11.01                  | 0                |
| 1                                                           | 0                         | 8.00                       | 0.70                   | 12.41                  |                  |
|                                                             |                           | 4.01                       | 0.22                   | 6.34                   |                  |
|                                                             |                           | 3.74                       | 0.24                   | 7.11                   | 0                |
| 1                                                           | 0                         | 5.10                       | 0.26                   | 7.41                   |                  |
|                                                             |                           | 8.02                       | 0.30                   | 7.23                   | 0                |
| 1                                                           | 0                         | 4.35                       | 0.32                   | 11.94                  | 0                |
|                                                             |                           | 4.45                       | 1.25                   | 22.16                  | 0                |
|                                                             |                           | 6.21                       | 1.14                   | 23.28                  | 0                |
| 10                                                          | 0                         | 4.10                       | 1.00                   | 14.04                  |                  |
|                                                             |                           | 3.01                       | 1.11                   | 21.52                  | ±                |
| 20                                                          | 0                         | 3.61                       | 1.29                   | 22.40                  |                  |
|                                                             |                           | 3.20                       | 1.28                   | 21.81                  | ±                |
| 24                                                          | 0                         | 0.70                       | 1.10                   | 19.02                  | ±                |
|                                                             |                           | 1.30                       | 1.21                   | 21.70                  | ±                |
| Dust Exposure Followed by Prolonged Residence in Normal Air |                           |                            |                        |                        |                  |
| 20                                                          | 3                         | 1.25                       | 1.41                   | 31.91                  |                  |
|                                                             |                           | 0.61                       | 1.16                   | 23.21                  | +                |
| 20                                                          | 6                         | 5.25                       | 0.66                   | 12.70                  | +                |
|                                                             |                           | 1.20                       | 0.67                   | 14.22                  | +                |
| 20                                                          | 12                        | 0.35                       | 0.91                   | 13.08                  |                  |
|                                                             |                           | 3.17                       | 0.61                   | 12.41                  | 1+               |

\*The symbols averaging the tissue reaction in each group represent merely the relative degree of reaction, ranging from 0 to ± (approximately) to 2+ (the maximum observed in this experiment). The relationship of activity only within this table and cannot be compared with symbols to other tables.

**Summary and Interpretation of Inhalation Experiment with 100 per Cent Ball-Milled Asbestos Dust.**—The tissue reaction observed in this experiment was not as intense as that in the previous investigation with short fiber asbestos. The reaction was slower in development and less extensive even though more dust accumulated in the lungs. Since there were fewer fibers longer than 3 microns in the material used in this experiment, the results tend to confirm the interpretation made in the summary of the previous short fiber experiment that the reaction is not primarily chemical in nature, and to support the impression that reduction in size of abestos fibers does not increase the biologic activity of asbestos inhaled into the lung.

The finding of long, asbestos fibers in animals that had inhaled ball milled material is an example of the difficulty of completely eliminating long fibers from a large volume of asbestos as required for an inhalation experiment.

In regard to the progression of the tissue reaction after the animals had been removed from the dust, observed in this experiment but not in the others, the following interpretation is offered: When the reaction is well developed at the termination of exposure, the contraction of the fibrous tissue obscures any progression that may have occurred; in this experiment, however, since the reaction observed was less mature, its subsequent progress was more readily apparent.

#### Long, Fine Asbestos Dust

Since inhalation of short fiber and of 100 per cent ball-milled asbestos dust did not result in acceleration of the tissue reaction in comparison with that produced by King's flints, the hypothesis that short fibers of asbestos were of minor importance in the etiology of asbestosis was given added support, and attention was directed to the view that the long fibers were of primary significance in that etiology. The King's flint asbestos used in the first inhalation experiment had a rather low content of fibrous chrysotile and contained considerable serpentine and other impurities. Therefore, it was decided to conduct a new inhalation experiment with a purer form of chrysotile which would be richer in long fibers.

**Composition and Amorphous Concentration of the Dust.**—The dusting material employed in this investigation was obtained from an asbestos fabricating plant. Samples of several varieties of long fiber asbestos dust were first submitted to the Saranac Laboratory for examination, and one of these, which was low in magnesian and fibrous content, estimated to be about 75 per cent, was selected as most suitable. Steel wire brushes, fastened to the inside surface of the hopper and to the rotating paddle as in the preceding inhalation experiment, were used to sweep up the bundles of asbestos and fibers into fibers into the atmosphere.

The composition of the long fiber asbestos used is indicated by the chemical and petrographic analysis given in tables 2 and 3. Analysis of air-suspended material from the dust room disclosed that about 20 per cent of the long fiber dust was chrysotile and about 20 per cent serpentine; as already noted, the composition of a similar air-throat sample of ball milled, short fiber dust was 15 per cent chrysotile and 60 per cent serpentine.

The dust concentration as revealed by unguetter samples taken inside the animal cages was much lower than the concentration for the experiments with short fiber or ball-milled dust. For the first year of the experiment with long fiber asbestos the average of the light field counts was 12 million particles per cubic foot of air; for the second year, 48 million; for the third year, 39 million; and for the fourth year, 43 million.

The size frequency of amorphous samples of the long fiber asbestos dust and of the ball milled dust is shown in table 11. Both samples were collected with the electrostatic precipitator. It will be noted that there was far more fibrous material in the long fiber dust.

Guinea pigs, rats and mice were employed in this inhalation experiment. The results, summarized in table 12, are described in greater detail below.

**Reaction in Guinea Pigs.**—The experiment was started with 100 guinea pigs. After exposure had been carried on for a year, a severe epidemic of pneumonia arose in the dust room and about one third of the animals died or were killed. To replace them, 38 more guinea pigs were added to the surviving group. Histological examination revealed lesions in the lungs after eight months of dust exposure, consisting of cellular connective tissue about the terminal bronchioles (fig. 7 A). At 12 months there were adenomatoid changes in the adjacent parenchymal areas, and by the sixteenth month (fig. 7 B) definite fibrosis was present in these areas, as well as around the bronchioles. The fibrous lesion could be seen macroscopically at 20 months. From this time on the reaction increased in extent and in the amount of collagen, and by the thirty-fourth month, it had fanned out

TABLE 11.—Size-Frequency of Amorphous Long Fiber and 100 per Cent Ball-Milled Asbestos Dust Collected Inside Cages

| Type of Asbestos | Guinea, % |         |      | Rats, % |         |      | Groups |
|------------------|-----------|---------|------|---------|---------|------|--------|
|                  | < 1       | 1 to 10 | > 10 | < 1     | 1 to 10 | > 10 |        |
| Long Fiber       | 61.4      | 31.1    | 0.0  | 33.4    | 67.1    | 19.0 | 100    |
| Ball milled      | 80.0      | 19.0    | 0.0  | 0.0     | 0.0     | 11.1 | 100    |

TABLE 12.—Summary of Inhalation Experiment with Long Fiber Asbestos Dust

| Nature of Exposure          | Animals         | Dust exposure concentration through out life | Maximum histological reaction after dust exposure |      | Results                                                                                    |
|-----------------------------|-----------------|----------------------------------------------|---------------------------------------------------|------|--------------------------------------------------------------------------------------------|
|                             |                 |                                              | Guinea pigs                                       | Rats |                                                                                            |
| Dust exposure for 10 months | 100 guinea pigs | 4 rats                                       | 20                                                | 0    | Definite fibrosis in 10 mo. slowly developing fibrosis first seen at 31 mo.                |
| Dust exposure for 12 months | 20 mice         | 11                                           | 0                                                 | 0    | Marked peribroncholar fibrosis first seen at 12 mo.                                        |
| Dust exposure for 16 months | 12 guinea pigs  | 20                                           | 14                                                | 0    | Clearing of inflammatory reaction and definite contraction of fibrous tissue               |
| Dust exposure for 20 months | 0 guinea pigs   | 27                                           | 0                                                 | 0    | Clearing of inflammatory reaction and slight contraction of fibrous tissue                 |
| Dust exposure for 34 months | 1 rat           | 11                                           | 11                                                | 11   | Similar to continuous exposure group; reaction of peribroncholar in one of the two animals |

considerably into the parenchyma (fig. 8 A). The lesions were rather sharply localized and the extensions from different bronchioles showed no tendency to fuse, even in animals exposed for the maximum period of three years. Although the intrapulmonary reaction sometimes reached the pleura, there was no involvement of that membrane. Emphysema was not detected at any point. Some thickening of the larger bronchi with a chronic inflammatory infiltration was revealed, but it was considered no more than would be produced by a similar period of inhalation of any dust.

In guinea pigs exposed to the dust for 20 months and then removed to normal air, there was a marked tendency for cellular inflammatory reaction to clear. This effect, accompanied by contraction of the fibrous tissue, resulted in a diminishing size of the focal lesions. None of these animals, killed at various periods up to 14 months after exposure, revealed lesions as large as those in the group killed at the end of the 20 month exposure period or those in animals which remained in the dust room for more than 20 months. Fourteen months after dust exposure ceased, the foci in four of the six remaining guinea pigs were so small that they were visible only with a hand lens (fig. 8 B).

begin to appear in the medulla, and by the fourteenth month most of the alveoli had been replaced by cellular connective tissue. This picture, which resembled that in early silicosis, persisted to the end of the experiment. Some animals showed, as a variant, heavy sheets of diffusely distributed monocytes and large active giant cells, but there was never any necrosis or hyaline formation. The spindle-shaped

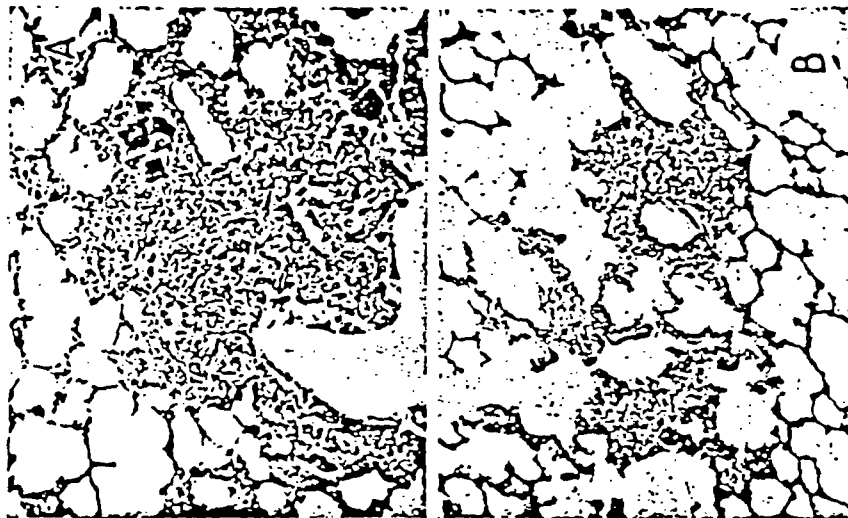


Fig. 8.—Long fiber asbestos inhalation experiment: A, lung of a guinea pig with 14 months' dust exposure. A bronchiole is seen at the lower center; the large area above it represents the involvement of alveolar walls. Compare with figure 7B and note the increased extent of reaction ( $\times 200$ ).

B, lung of a guinea pig with 20 months' dust exposure and then 14 months' living in normal air. The reaction is essentially like that shown in figure 7B. The bronchiole at the right center is surrounded by fibrous tissue with alveolar change at the right. There is residual scarring in the walls of adjacent alveoli at the left. It is apparent that no progression has occurred ( $\times 200$ ).

new cells were yellowish from fine pigment granules that stained for iron. No fibers or asbestos bodies were seen.

In the groups exposed for 27 months, and then transferred to a normal atmosphere the response was quite similar to that in the 20 month exposure animals mentioned above. Small but well always visible on gross inspection of sections of all guinea pigs of the 27 month series, but in no instance was there evidence of the reaction.



Fig. 7.—Long fiber asbestos inhalation experiment: A, lung of a guinea pig with eight months' dust exposure. The bronchiole at the center already shows an accumulation of phagocytes, cells, and there is a slight deposition of collagen. Compare with figure 6B, showing the reaction in half milled asbestos after 24 months' exposure ( $\times 200$ ).

B, lung of a guinea pig with 16 months' dust exposure. Again note a bronchiole with its surrounding reaction, consisting of fibrosis and alveolar change. Collagen deposition is now seen in the walls of adjacent alveoli at the right ( $\times 200$ ).

In the tracheobronchial lymph nodes reaction was first visible at the third month of exposure. By the eighth month patches of cellular connective tissue

Although asbestos bodies were found in the lung as early as one month after exposure, before they were rare and hard to find. At five months more were visible, chiefly coated macrophages, and at eight months many bodies

TABLE 13.—Analysis of Lung of Guinea Pig Exposed to Dust in Inhalation Experiment with Long Term (10-15) Rat Dust

| Exposure to Dust, Mo. | Period in Normal Air, Mo. | And of Air Contained in Lung | Total Dust Exposed to Guinea Pig | Total No. of Guinea Pigs | Tumor Reaction* |
|-----------------------|---------------------------|------------------------------|----------------------------------|--------------------------|-----------------|
| 1                     | 0                         | 4.3                          | 0.04                             | 10                       |                 |
|                       |                           | 4.4                          | 0.04                             | 11                       |                 |
|                       |                           | 4.5                          | 0.04                             | 9                        |                 |
| 3                     | 0                         | 4.6                          | 0.05                             | 12                       |                 |
|                       |                           | 4.7                          | 0.05                             | 13                       |                 |
|                       |                           | 4.8                          | 0.04                             | 14                       |                 |
| 5                     | 0                         | 4.9                          | 0.05                             | 15                       |                 |
|                       |                           | 5.0                          | 0.04                             | 16                       |                 |
|                       |                           | 5.1                          | 0.04                             | 17                       |                 |
| 8                     | 0                         | 5.2                          | 0.04                             | 18                       |                 |
|                       |                           | 5.3                          | 0.04                             | 19                       |                 |
|                       |                           | 5.4                          | 0.04                             | 20                       |                 |
| 10                    | 0                         | 5.5                          | 0.04                             | 21                       |                 |
|                       |                           | 5.6                          | 0.04                             | 22                       |                 |
| 12                    | 0                         | 5.7                          | 0.04                             | 23                       |                 |
|                       |                           | 5.8                          | 0.04                             | 24                       |                 |
| 14                    | 0                         | 5.9                          | 0.04                             | 25                       |                 |
|                       |                           | 6.0                          | 0.04                             | 26                       |                 |
| 16                    | 0                         | 6.1                          | 0.04                             | 27                       |                 |
|                       |                           | 6.2                          | 0.04                             | 28                       |                 |
| 18                    | 0                         | 6.3                          | 0.04                             | 29                       |                 |
|                       |                           | 6.4                          | 0.04                             | 30                       |                 |
| 20                    | 0                         | 6.5                          | 0.04                             | 31                       |                 |
|                       |                           | 6.6                          | 0.04                             | 32                       |                 |
| 22                    | 0                         | 6.7                          | 0.04                             | 33                       |                 |
|                       |                           | 6.8                          | 0.04                             | 34                       |                 |
| 24                    | 0                         | 6.9                          | 0.04                             | 35                       |                 |
|                       |                           | 7.0                          | 0.04                             | 36                       |                 |
| 26                    | 0                         | 7.1                          | 0.04                             | 37                       |                 |
|                       |                           | 7.2                          | 0.04                             | 38                       |                 |
| 28                    | 0                         | 7.3                          | 0.04                             | 39                       |                 |
|                       |                           | 7.4                          | 0.04                             | 40                       |                 |
| 30                    | 0                         | 7.5                          | 0.04                             | 41                       |                 |
|                       |                           | 7.6                          | 0.04                             | 42                       |                 |
| 32                    | 0                         | 7.7                          | 0.04                             | 43                       |                 |
|                       |                           | 7.8                          | 0.04                             | 44                       |                 |
| 34                    | 0                         | 7.9                          | 0.04                             | 45                       |                 |
|                       |                           | 8.0                          | 0.04                             | 46                       |                 |
| 36                    | 0                         | 8.1                          | 0.04                             | 47                       |                 |
|                       |                           | 8.2                          | 0.04                             | 48                       |                 |
| 38                    | 0                         | 8.3                          | 0.04                             | 49                       |                 |
|                       |                           | 8.4                          | 0.04                             | 50                       |                 |
| 40                    | 0                         | 8.5                          | 0.04                             | 51                       |                 |
|                       |                           | 8.6                          | 0.04                             | 52                       |                 |
| 42                    | 0                         | 8.7                          | 0.04                             | 53                       |                 |
|                       |                           | 8.8                          | 0.04                             | 54                       |                 |
| 44                    | 0                         | 8.9                          | 0.04                             | 55                       |                 |
|                       |                           | 9.0                          | 0.04                             | 56                       |                 |
| 46                    | 0                         | 9.1                          | 0.04                             | 57                       |                 |
|                       |                           | 9.2                          | 0.04                             | 58                       |                 |
| 48                    | 0                         | 9.3                          | 0.04                             | 59                       |                 |
|                       |                           | 9.4                          | 0.04                             | 60                       |                 |
| 50                    | 0                         | 9.5                          | 0.04                             | 61                       |                 |
|                       |                           | 9.6                          | 0.04                             | 62                       |                 |
| 52                    | 0                         | 9.7                          | 0.04                             | 63                       |                 |
|                       |                           | 9.8                          | 0.04                             | 64                       |                 |
| 54                    | 0                         | 9.9                          | 0.04                             | 65                       |                 |
|                       |                           | 10.0                         | 0.04                             | 66                       |                 |
| 56                    | 0                         | 10.1                         | 0.04                             | 67                       |                 |
|                       |                           | 10.2                         | 0.04                             | 68                       |                 |
| 58                    | 0                         | 10.3                         | 0.04                             | 69                       |                 |
|                       |                           | 10.4                         | 0.04                             | 70                       |                 |
| 60                    | 0                         | 10.5                         | 0.04                             | 71                       |                 |
|                       |                           | 10.6                         | 0.04                             | 72                       |                 |
| 62                    | 0                         | 10.7                         | 0.04                             | 73                       |                 |
|                       |                           | 10.8                         | 0.04                             | 74                       |                 |
| 64                    | 0                         | 10.9                         | 0.04                             | 75                       |                 |
|                       |                           | 11.0                         | 0.04                             | 76                       |                 |
| 66                    | 0                         | 11.1                         | 0.04                             | 77                       |                 |
|                       |                           | 11.2                         | 0.04                             | 78                       |                 |
| 68                    | 0                         | 11.3                         | 0.04                             | 79                       |                 |
|                       |                           | 11.4                         | 0.04                             | 80                       |                 |
| 70                    | 0                         | 11.5                         | 0.04                             | 81                       |                 |
|                       |                           | 11.6                         | 0.04                             | 82                       |                 |
| 72                    | 0                         | 11.7                         | 0.04                             | 83                       |                 |
|                       |                           | 11.8                         | 0.04                             | 84                       |                 |
| 74                    | 0                         | 11.9                         | 0.04                             | 85                       |                 |
|                       |                           | 12.0                         | 0.04                             | 86                       |                 |
| 76                    | 0                         | 12.1                         | 0.04                             | 87                       |                 |
|                       |                           | 12.2                         | 0.04                             | 88                       |                 |
| 78                    | 0                         | 12.3                         | 0.04                             | 89                       |                 |
|                       |                           | 12.4                         | 0.04                             | 90                       |                 |
| 80                    | 0                         | 12.5                         | 0.04                             | 91                       |                 |
|                       |                           | 12.6                         | 0.04                             | 92                       |                 |
| 82                    | 0                         | 12.7                         | 0.04                             | 93                       |                 |
|                       |                           | 12.8                         | 0.04                             | 94                       |                 |
| 84                    | 0                         | 12.9                         | 0.04                             | 95                       |                 |
|                       |                           | 13.0                         | 0.04                             | 96                       |                 |
| 86                    | 0                         | 13.1                         | 0.04                             | 97                       |                 |
|                       |                           | 13.2                         | 0.04                             | 98                       |                 |
| 88                    | 0                         | 13.3                         | 0.04                             | 99                       |                 |
|                       |                           | 13.4                         | 0.04                             | 100                      |                 |

\* The tumor reaction in each group represents merely the tumor reaction observed from the lungs of the animals in the group. The tumor reaction in the lungs of the animals in the group is compared with the tumor reaction in the lungs of the animals in the group.

were free in connective tissue. They became fairly abundant as exposure continued, although in some later animals the asbestos bodies were only moderately numerous.

It is important to note from analyses of the lungs (table 14) that even though the tissue decayed at any given period of time was much greater in the guinea

pigs of this experiment than in those exposed to either short fiber or ball-milled asbestos, the amount of mineral matter in the lung ash was much less.

**Reaction in Cats.**—Four cats inhaled the long fiber asbestos dust for periods of 14, 25, 33 and 42 months, respectively, and were immediately killed. Two other cats, after being exposed to dust for 18 months, lived in a normal atmosphere for an additional 24 months. Fourteen months' exposure was sufficient to produce cellular accumulations of phagocytes around terminal bronchioles and peripheral arterioles together with compact collections of similar cells in the tracheobronchial lymph nodes. At that time there were no typical asbestos bodies, but smooth, pointed, yellow fibers were seen very rarely. With continued exposure, up to 42 months, reaction in the lungs noted progressed to the formation of cellular connective tissue which made well defined sheaths about the respiratory bronchioles and arterioles, marked lymphoid hyperplasia and lymphoid infiltration of bronchiolar walls (fig. 9). Typical asbestos bodies were not formed, although there was



Fig. 9.—Long fiber asbestos inhalation experiment: Lung of a cat with 42 months' dust exposure. Two bronchioles are shown with adjacent cellular reaction and collagen deposition (X 200).

an occasional fiber, smooth, yellow and pointed. Pleurisy was not present. The reaction was similar in location to that in the guinea pigs, but fibrosis was much slower in development. Roentgenograms of cats made after exposure periods of 25, 33 and 42 months, respectively, failed to demonstrate evidence of pulmonary lesions.

**Reaction in Rats.**—Although 20 rats were placed in the dust room, many died from pneumonia and were not suitable for study. Five animals, of which one was exposed for 19 months and four for 25 months, were free from pulmonary infection and offered a basis for tentative conclusions. In the 19 month animal, the reaction was just beginning. All four animals killed at 25 months showed a well marked peribronchiolar fibrosis. After a lung search, only two small, smooth asbestos bodies were found in the 19 month animal and none was found in the 25 month animal. Thus these animals exhibited fibrosis without asbestos bodies or fibrosis accompanied by only a very infrequent asbestos body.

TABLE 14.—Comparison of Reactions to Chrysole and Serpentine Injected Intratracheally

Dosage: Each animal was given an intratracheal injection of a 2 per cent suspension of the dust. Two weeks later another injection was given. Total amount of dust injected was 50 mg.

Animals used). Six groups of 6 guinea pigs each (one group for each type of dust). One or two animals in each group at 1, 2, 4, 6, 8 and 12 months after last injection.

incubation at 30°C. for 24 hr., dried and reprecipitated of dust; Chrysotile (best milled) untreated: Don milled for 1.76 hr., dried and reprecipitated in acetate buffer.

ground in acetone buffer 2 or 3 min.

*Chrysosoma (Chrysosoma) unicolor*: Grouped in eagle mortar to pass 500 mesh. Subweb material tested by J. Br. at about 100 C. No further

granulation.  
Ball milled for 1,736 hr. dried and reground in agate mortar.  
(ball milled) ignited: Ball milled acetylene treated for 8 hr. at about 700 C., then reground in agate mortar 2 or 3 min.

| Microfl.                                | Size of<br>Dust Inhibits<br>less | Results                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          |
|-----------------------------------------|----------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Crescentic<br>(bail inhib.)<br>undusted | 3 microns and<br>less            | Girding destroyed capacity to cause fibrosis. At 1 mo. resulting inflammation of air and cellular proliferation and contraction favored subnormal expansion of air spaces given off directly from terminal bronchioles. Reaction area became smaller with progress of time; no new fibrosis involved. No fibrous grayish area at periphery extending immediately adjacent. At 3 mo considerable inflammatory edema and loss of cellular proliferation. At 5 mo. with marked cellular proliferation and contraction resulting in small terminal bronchioles. This reaction occurred behind remaining bronchioles had limited extension of alveolar tissue. At 8 mo. reaction less extensive than at 5 mo. Apparently due to contraction of fibrous tissue; alveolar bodies were abundant. At 9 mo., reaction still less extensive, confined to the immediate vicinity of the small terminal bronchioles, where the air tissue was quite dense and was becoming fibrous in character. Sometimes it even infiltrated the bronchioles. At 10 mo., the air-tissue bodies had become scarce. At 11 mo., the well developed peribronchial and interbronchial alveolar areas of fibrous body produced considerably the same reaction. More peripherally are patches of pneumonitis with eosinophilic infiltration, some of which was later transformed into fibrous tissue. This seemed to be precursor of the localized, diffuse patches of false alveolar wall fibrosis and emphysema. |
| Chrysoidia<br>(bail inhib.)<br>lighted  | 20-20 microns<br>approx.         | Reaction limited to large forerun body giant cells without proliferation. Infiltrating the alveoli, which made firm brittle, destroyed their capacity to produce significant reaction.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| Serpentine<br>(bail inhib.)<br>undusted | 3 microns and<br>less            | Dust relatively inactive. At 1 and 2 mo. simple phagocytic possibly lymphoid cell infiltration; at 4 mo. a slight chronic pneumonitis; at 12 mo. only a little pneumonitis without augmentation of fibrosis.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     |
| Serpentine<br>(bail inhib.)<br>lighted  | 3 microns and<br>less            | Dust relatively inactive. Reaction essentially the same as for undusted serpentine. With lighted serpentine, less tendency for dust to be carried to bronchial pulps.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |

*Reaction to Ure.*—That of 20 chicks was used in this experiment. Within a year after most of them had died or were killed without showing an appreciable degree of pulmonary infection, the reaction to the inhaled dust was limited to phagocytes by macrophage cells. Usually these were widely scattered through the air spaces, a limited number were grouped about the terminal bronchioles, producing a zone of thickening of that wall. There was no suggestion of fibrosis.

Since then, adolestary leeches were observed in animals killed late in the experiment. Thus these animals, excluded of adolestary leeches without obvious

**Summary and Interpretation of Inhalation Experiment with Long Fiber Asbestos Dust.**—The purpose of this experiment was to evaluate the importance of long fibers in the tissue response to inhaled asbestos. The results, in comparison with those of previous investigations, indicate strongly that long fibers are chiefly responsible for asbestosis. Thus, the reaction in guinea pigs developed earlier and became more extensive in this experiment than in previous experiments in spite of a smaller concentration of amorphous dust and a lower mineral content of the lungs. Furthermore, typical peribronchovascular fibrosis was produced in alveoli, although in a previous experiment with short fiber dust peribronchovascular fibrosis did not develop in this species.

The cause of the cellular fibrosis in the lymph nodes of the guinea-pigs is not clear. It did not occur in other inhalation experiments with asbestos.

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Since the inhalation experiments reported above strongly suggested that long fibers of asbestos are the significant factor in the causation of a disease, a series of injection experiments was inaugurated wherein the dosage and the length of the fibers could be controlled more precisely.

Also, by the use of controlled dosages, the relative capacities of various excretory minerals to produce reaction could be compared. In these comparison experiments, guinea pigs, rabbits, rats and dogs were used, and the mineral diet was injected by the intratracheal, the intraperitoneal and the intravenous technique, but not all the techniques were used for each species. For the purpose of simplification the findings in each series of tests, except for dogs, have been condensed and reported in tables, to which reference will be made later. In the case of dogs, only one test was made, and since the findings were negative, no detailed report is included.

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As the asbestos minerals do not cause typical advanced fibrosis in extrapulmonary tissue, the intratracheal technique is the preferred way of introducing fibrogen dust into the experimental animal. In this method the dust suspension is injected by means of a special needle or catheter deep into the trachea, from which it flows into the lungs.

*Comparison of Fibrous and Nonfibrous Parts*—To demonstrate that the ability of adhesives to produce adheres relies on its adhesion character, the series of structural elements reported in Table 11 were performed

The tests were made with long fiber chrysotile, untreated, and with chrysotile that had been ignited to destroy its flexible structure or ball milled to reduce the length of fiber to 3 microns and less. At the same time control tests were made with serpentine, which has the same chemical composition as chrysotile but is nonfibrous. A review of the findings reveals that only the untreated, long fiber chrysotile produced typical peribronchiolar fibrosis and that ball-milled material containing only fibers less than 3 microns in length failed to cause fibrosis (figs. 10 and 11). Fibers subjected to ignition also had lost their capacity to cause serious tissue damage. Ignition produced important changes in the chrysotile fibers, among them being loss of water, an alteration from a flexible to a brittle structure and possibly other changes. Experi-



Fig. 11.—Serpentine injection experiment: Lung of a Guinea pig that had received an intratracheal injection of this dust four months before. A bronchiole is shown at the left center. The phagocytic cells exhibit little deflection for the bronchiole and collagen deposition is absent (X 200).

mental studies concerning this observation will be reported in a separate publication.

**Comparison of Various Long Fiber Dusts.**—Some very interesting findings are disclosed by the results of the experiments recorded in table 15. First, all the long fiber asbestos minerals tested, with the exception of anthophyllite, produced typical fibrosis. The characteristic peribronchiolar reaction caused by three representative long fiber asbestos minerals—chrysotile, amosite and crocidolite—is shown in figures 10, 11 and 12. Why anthophyllite behaved differently from the other asbestos minerals is not entirely clear.

Second, with the mineral brucite, which is not a silicate but is a fibrous form of magnesium hydroxide, a characteristic fibrosis like

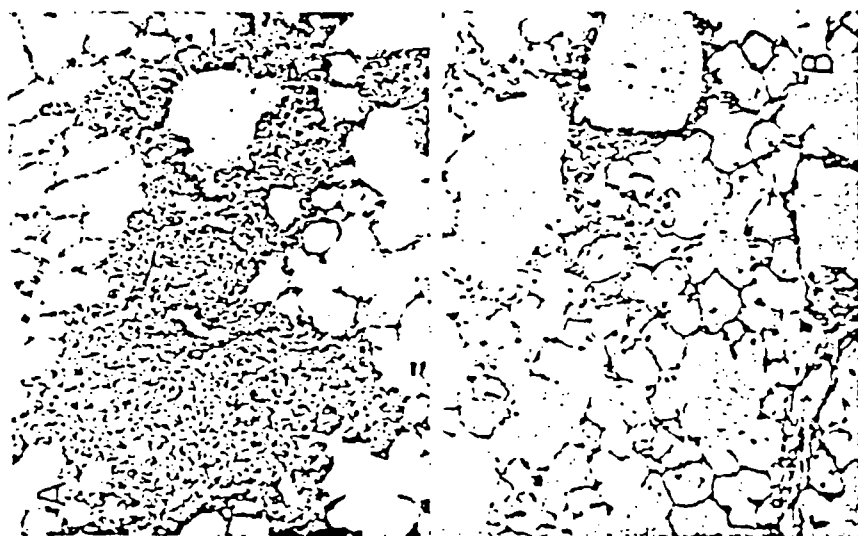


Fig. 10.—Comparison of reactions provoked by injected long fiber and ball-milled asbestos dusts: A lung of a Guinea pig which four months before had received an intratracheal injection of long fiber asbestos dust. Note the peribronchiolar accumulation of cells with collagen deposition. The bronchiole chiefly involved is in the middle of the reaction (X 200).

B, lung of a Guinea pig which four months before had received an intratracheal injection of ball-milled asbestos dust. A bronchiole is shown at the right. In contrast with A, note that only a few cells have accumulated about the bronchiole and that collagen deposition is absent (X 200).

that produced by the asbestos minerals was obtained (fig. 13 *d*). Since the brucite used contained only 0.90 per cent silica as an impurity, it is obvious that a siliceous component is not an essential factor in the development of asbestosis.



Fig. 12.—Anostic and crocidolite injection experiments: *A*, lung of a guinea pig four months after an intratracheal injection of anostic. The inflammatory reaction exhibits pronounced accumulation of cells and collagen deposition ( $\times 200$ ). *B*, lung of a guinea pig four months after an intratracheal injection of crocidolite. As in *A*, peribronchovascular accumulation of cells and deposition of collagen are shown ( $\times 200$ ).

Third, no fibrosis resulted from the injection of glass wool fibers (fig. 13B), even though glass wool resembles asbestos in many ways. However, there are fundamental differences. A glass wool fiber 3 microns

foregoing. Their objective is to use the effect of a high seed population given two months after sowing to increase the yield of the crop.

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At 150 sec after infection, erythrocytes, leukocytes, and platelets were found in the lymphatic nodes. At 1 min, early erythrocytopenia, leukopenia, and thrombocytopenia were observed. Lymphocyte function at 6 hr, 2 weeks after infection was similar to that of lymphocytes from the lymphatic nodes of uninfected mice. *Stress of lymphoblast growth* in mouse lymphatic nodes was observed at 1 min, *stress of lymphoblast growth* in mouse lymphatic nodes was observed at 1 min, *stress of lymphoblast growth* in mouse lymphatic nodes was observed at 1 min.

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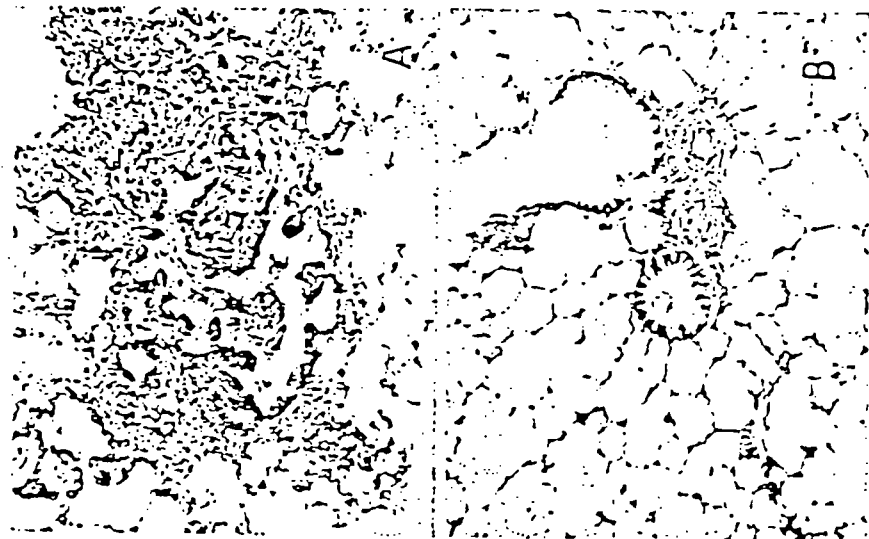


Fig. 13—Brucite and glass wool infection experiment. *A*, lining of a Kuma-  
yama which four months before had received an oral attached infection of brucite.  
Even with this inoculum no bacteria could there be produced in the animal tissue  
and cells and disposition of collagen similar to that shown in *A* and *B* of figure 12  
(*cf.* text).

*B.* Ring of a femur for which four months before had received an intramedullary cast, there is a glazy wall. Two lamellae are shown, one in cross section and the other in longitudinal section. Below the latter is a thick walled blood vessel. The lamellae are without traction and can be considered normal for comparison with other figures. Glazy wall fibers are present in this field but cannot be seen at higher magnification (Fig. 29).

TABLE 16.—Comparison of Reactions Produced by Long Fiber and Short Fiber Darts Injected Intratracheally

**Dosage:** Two injections of 0.5 cc. of a 5 per cent suspension given two weeks apart. Total volume was 1.0 cc.

| Microal                   | Blue of<br>Dust Fatigue                                                  | Results                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               |
|---------------------------|--------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Chromella<br>(Thaloid)    | Long fiber,<br>20-30 microns<br>Short fiber,<br>5-10 microns<br>and less | A distinct fibrous matter to chromella (fibrous) un-<br>breathed in table II.<br>No reaction to ferruginous (dust mixed) unbreathed<br>in table II.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   |
| Amurella                  | Long fiber,<br>20-30 microns<br>Short fiber,<br>20 microns<br>and less   | Typical fibrous embolobronchiolitis and peribronchiolitis.<br>Refer to table I3.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      |
|                           |                                                                          | Reaction limited to phagocytosis with lymphocytic infil-<br>tration of alveolar walls. Short fibers packed inside<br>of alveoli, but not outside. Some fibers, some coated in<br>form of thin, translucent, wavy lines, some in form of<br>long, thread-like, wavy lines. Some fibers, some coated<br>in liquid, alveoli contained fluid and small cells; most<br>phagocytes were within air spaces and had not<br>migrated in walls. At 4 mo., few attachment dots<br>had worked themselves into interstitial tissue, where<br>they were minute proliferations of lymphoid cells and<br>macrophages. Some of the latter were in contact with<br>reaction with some pneumonitis, no bronchiolitis<br>Typical alveolar lumina present.                                                                 |
| Cricoballia<br>(Bullaria) | Long fiber,<br>20-30 microns<br>Short fiber,<br>20 microns<br>and less   | Alveolated fibrous embolobronchiolitis and peribronchio-<br>litis. Refer to table I3.<br>No reaction to table I3.<br>No reaction to table I3.<br>Walls heavily infiltrated with monocytes and lympho-<br>cytic cells. At 4 mo., a moderate degree of epithelial<br>induration of walls, small giant cells packed with<br>dust spicules. At 6 and 84 mo., masses of giant cells,<br>containing mineral particles, in small bronchi but not<br>in alveoli. At 84 mo., some giant cells were in-<br>fected by terminal air spaces. Numerous alveolar<br>bodies. No reaction to connective tissue. No colo-<br>bronchial proliferation. At 13 mo., many scattered<br>small monocytes packed with dust. No reduplica-<br>tions. No peripheral fibrosis. In lymph node, slight<br>lymphocytic infiltration. |
| Anthrophylla              | Long fiber,<br>20-30 microns<br>Short fiber,<br>20 microns<br>and less   | Emphysematous bronchiolitis and giant cells but no definite<br>fibrosis. Refer to table I3.<br>No fibrosis and practically no alveolar bodies. At 1<br>mo., local collections of dust-filled monocytes and a<br>few giant cells; at 1 mo., some alveolar epithelial<br>reaction; at 2 mo., minute pneumonitis with mono-<br>cytic infiltration of alveolar walls. At 4 mo., some<br>isolated collections of dust cells inside air spaces<br>about terminal arterioles. Reaction in walls limited<br>to lymphoid cell infiltration. No fibrosis. In lymph<br>node, reaction limited to slight prominence of lym-<br>phocytes about bronchioles. Refer to table I3.                                                                                                                                     |
| Trypanella                | Long fiber,<br>20-30 microns<br>Short fiber,<br>20 microns<br>and less   | Simple foreign body reaction. No acute inflammation.<br>No accumulation of dust at or about terminal lung<br>arteries. No embolobronchiolitis. At 1 mo., scattered small<br>giant cells in alveoli. At 2 mo., some alveolar walls<br>with monocytes and lymphoid cells. At 1 mo.<br>little change except more cellular infiltration of co-<br>nective tissue. At 4 mo., lymphoid infiltration and<br>thinning of walls about some but not all terminal<br>bronchioles.                                                                                                                                                                                                                                                                                                                                |
| Buclia                    | Long fiber,<br>20-30 microns<br>Short fiber,<br>20 microns<br>and less   | Typical embolobronchiolitis and peribronchiolitis<br>like reaction to asbestos minerals. Refer to table I3<br>first type of reaction. At 1 mo. after injection, small<br>monocytes wholly retained through air spaces; focus<br>of asbestos with lymphoid infiltration of connective<br>tissue walls. No embolobronchial reaction at, with<br>only a few alveolar cells. At 2 mo., some alveolar<br>walls typical alveolar infiltration. At 4 mo., small<br>clumps of inactive dust filled phagocytes; no fibrosis.<br>No reaction to lymph nodes.                                                                                                                                                                                                                                                    |

*Comparison of Long Fiber and Short Fiber Dusts.*—With quartz dust it has been demonstrated that the smaller the particles the more

minerals is the same reaction and that particles larger than 3 microns in diameter cause little reaction. In the case of asbestos, however, the reverse is true and apparently only long fibers have any specific effect, as was suggested by the inhalation experiments. This is confirmed by the data of table IV, in which a series of tests with fibrous minerals is reported. When the injected dust consisted of fibers 20 to 50 microns long, all the fibrous minerals tested except anthophyllite, as noted in the preceding section, produced fibrosis; when the material was prepared by first grinding the fibrous dust until the length of fibers was reduced to 20 microns and less or, in some cases, to 3 microns and less, none of the injected dusts caused fibrosis.

These results differ from those of King, Clegg and Rae,<sup>14</sup> who represented the production of reticulation comparable to the experimental fibrotic nodules in rabbits receiving monthly intratracheal injections of 100 mg of Khlofasin asbestos fibers, 15 microns long, and the production of diffuse interstitial fibrosis in rabbits receiving similar injections of short fibers, 2.5 microns in length. We believe this dose, especially in the long term rabbits, is highly excessive. In our experiments the dosage was kept low in order to minimize untoward reactions which might obscure the peritoneal nodular type of fibrosis which characterizes naturally occurring asbestososis.

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The experiments summarized in table 17, in which the intravenous method of injection was employed, show that the asbestos minerals are far different from quartz in their action on tissue. It has been repeatedly demonstrated that intravenous injection of quartz particles 3 microns and less in diameter will cause a typical tissue reaction with the development of a fibrotic lesion, in extrapulmonary sites, such as the liver and the spleen. Asbestos minerals, however, on intravenous injection generally produce only an inert type of reaction, as is revealed by the results given in the table. The reason for the early deaths in the experiment with chrysotile particles is not clear.

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The results of injection experiments with the infraperitoneal to lung route are given in table 16. It will be noted that the lung fibrocyte division produced a fibrous reaction while ducts composed of particles 3 microns and less in size caused only an inert type of response. These experiments indicate also that the fibrosis initiated by the irritation of asbestos fibers is not restricted to the lungs, as was formerly assumed, but can be produced in the testis, as well.

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A number of additional experiments were conducted to throw more light on specific phases of the advection problem.

H. King, F. J. Chapp, J. W. and R. G. V. M. Effect of Abores, and of Abores, on the Absorption of Light, *Trans. L.I.S. 1940, 4, 10-11.*

TABLE 17.—Summary of Injection Experiments by Intraprostatic Technique

When colloidal aluminum hydroxide had been added to a suspension of long fiber chrysotile prior to injecting this suspension intratracheally into rats, the aluminum compound did not prevent the irritation of tissue due to chrysotile. If anything, the acute inflammatory response evoked by the injected fibrous mineral was accelerated. One month after the last injection of the dust suspension the bronchiolitis was becoming fibrous. King and his associates also found that aluminum failed to protect pulmonary tissue from the irritation caused by asbestos fibers<sup>11</sup>; in their experiments metallic aluminum was used instead of the hydroxide.

#### FORMATION OF ASBESTOS BODIES

The iron in the coating of the asbestos body appears to be derived from blood or tissue elements and not, as has been suggested, from the mineral fiber. After two kinds of chrysotile were injected subcutaneously into the groin of a guinea pig—one kind containing 2 per cent and the other 0.2 per cent ferric oxide—the asbestos bodies were equally numerous at both sites of injection and showed no difference in their reaction to prussian blue, the reagent which stains iron. This finding is in agreement with that of Giroux.<sup>12</sup>

#### TISSUE REACTION TO ASBESTOS BODIES

Asbestos bodies recovered from human lung tissue and injected intratracheally into guinea pigs failed to produce a fibrous reaction. The material for injection was obtained by digesting with sodium hypochlorite solution the lung tissue removed at autopsy from an asbestos worker. The asbestos bodies could be seen in the guinea pigs for at least a year after injection. This experiment shows that the asbestos body has a rather resistant coating which is not destroyed by moderate hypochlorite treatment, which may be maintained in vivo for a year or longer and which renders the fiber incapable of producing fibrosis. It thus appears that the coating is a protective mechanism. This thought was expressed by Heimker as early as 1934.<sup>13</sup>

#### THEORY OF IRRITANT ACTION

Two hypotheses have been proposed to explain the tissue irritation and reaction caused by asbestos fibers: the chemical and the mechanical. In the chemical theory, which is based on experience with quartz, it is assumed that the asbestos minerals dissolve in the body fluids and that in this process their bases are leached away to leave silica in a form capable of irritating tissues. According to this hypothesis asbestosis is merely an indirect silicosis. Several facts make the chemical theory untenable: Intratracheal injection of brittle fibers, which had a silica

12. Giroux, M.: *Amiantose expérimentale: valeur pathogénomique du "corps d'amianté"*. Laval méd. 8:239, 1943.

13. Heimker, E.: *Über die Asbestkörperchen: Bemerkungen zu der Arbeit von Rieger*. Virchows Arch. f. path. Anat. 293:527, 1934.

TABLE 15.—Summary of Injection Experiments by Intraperitoneal Technique

| Experiment | Material              | Amount of dust injected                 | Time of observation | Reaction    |
|------------|-----------------------|-----------------------------------------|---------------------|-------------|
| 1          | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 2          | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 3          | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 4          | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 5          | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 6          | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 7          | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 8          | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 9          | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 10         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 11         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 12         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 13         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 14         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 15         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 16         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 17         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 18         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 19         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 20         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 21         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 22         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 23         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 24         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 25         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 26         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 27         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 28         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 29         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 30         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 31         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 32         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 33         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 34         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 35         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 36         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 37         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 38         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 39         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 40         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 41         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 42         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 43         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 44         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 45         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 46         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 47         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 48         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 49         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 50         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 51         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 52         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 53         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 54         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 55         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 56         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 57         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 58         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 59         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 60         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 61         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 62         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 63         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 64         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 65         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 66         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 67         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 68         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 69         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 70         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 71         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 72         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 73         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 74         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 75         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 76         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 77         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 78         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 79         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 80         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 81         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 82         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 83         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 84         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 85         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 86         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 87         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 88         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 89         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 90         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 91         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 92         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 93         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 94         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 95         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 96         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 97         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 98         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 99         | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |
| 100        | Long fiber chrysotile | 0.1 cc. of 0.5 per cent dust suspension | 10 days             | No reaction |

content of only 0.4 per cent. caused typical fibrosis like that produced by the asbestos minerals; free silica particles increase in potency as the particle size becomes less, but asbestos fibers shorter than about 10 to 20 microns are relatively innocuous; aluminum hydroxide molecules the irritating effect of quartz but not of asbestos; serpentine has the same chemical composition as flog, talc, clay, etc., but it produced only an inert type of tissue reaction; there is a wide range in the chemical composition of the minerals which do cause asbestosis (table 10). In view of this evidence it seems more likely that asbestosis is caused by an unusual mechanical irritation due to long asbestos fibers, this irritation being related to the peculiar filamentary structure of the fiber and the associated flexibility, which are possessed by no other foreign body studied. Thus, ignition of cigarette filters changed their structure and made them inert, although the same fibers, before being heated, would have produced fibrosis (table 11). Further support for the theory of mechanical irritation is that asbestosis occurs in an organ of high mobility—the lung—and that a fibrous reaction can be produced by injecting

TABLE 10. *Analysis of Various Minerals*

| Mineral            | SiO <sub>2</sub> | FeO  | Al <sub>2</sub> O <sub>3</sub> | CaO | MgO  | Na <sub>2</sub> O | K <sub>2</sub> O | Loss | Total |
|--------------------|------------------|------|--------------------------------|-----|------|-------------------|------------------|------|-------|
| Asbestos           | 46.2             | 1.0  | 11.5                           | 1.0 | 3.0  | 4.4               | 0.1              | 0.9  | 67.1  |
| Amphibole          | 52.4             | 1.0  | 1.2                            | 1.2 | 12.2 | 21.7              | 0.1              | 0.2  | 89.8  |
| Aluminum hydroxide | 38.7             | 0.2  | 0.2                            | 0.4 | 31.3 | 0.1               | 0.1              | 0.2  | 69.0  |
| Quartz             | 99.9             | 0.1  | 0.0                            | 0.0 | 0.0  | 0.0               | 0.0              | 0.0  | 100.0 |
| Chrysotile         | 35.2             | 1.1  | 0.1                            | 0.0 | 34.3 | 0.0               | 0.0              | 0.0  | 70.6  |
| Chrysotile         | 34.9             | 10.2 | 4.7                            | 1.0 | 12.3 | 0.0               | 0.0              | 0.0  | 63.1  |
| Tremolite          | 34.9             | 1.1  | 0.4                            | 0.0 | 30.2 | 0.0               | 0.0              | 0.0  | 66.6  |

asbestos fibers into the peritoneum, where there is also a degree of mobility, but not by injecting them into other extrapulmonary organs such as the liver, the spleen and subcutaneous tissue.

#### CONCLUSIONS

The experimental investigations with asbestos minerals were confined primarily with the effect of the dust on natural tissue, but some attention was given to other phases, such as susceptibility to infection. The only experiment in which the effect of asbestos dust on a pulmonary infection was studied was the first inhalation experiment, carried on with King's death dust. It is unfortunate that, owing to the lack of adequate facilities at that time, infection studies could not be made in the other inhalation experiments also.

#### SUSCEPTIBILITY TO TUBERCULOUS INFECTION

The development of a tuberculous process initiated at the beginning of exposure to dust, and also of a tuberculous infection superimposed on an established asbestosis was described in preceding sections of this paper. It may be stated that asbestos when classified according to the effect of a dust on tuberculous infection would be placed below an active

dust like quartz but above an inert dust such as iron oxide. In animals infected with attenuated tubercle bacilli, quartz causes the infectious process to progress until the animal dies of tuberculosis. Inert dusts have no effect on the infection, and the lesions usually heal and the disease disappears. Asbestos dust is in a different category. In the experimental investigation, when the fibrous dust was being inhaled during the evolution of the infection, there was spreading of the tuberculous process for a time, but usually the stimulus for continued proliferation of the tubercle bacilli was not sustained, the progression was arrested and healing followed. In guinea pigs injected with attenuated tubercle bacilli after being exposed to asbestos dust for slightly more than two years, progressive disease did not develop. The only modification of the infection was one of localization, a few bacilli being retained in the fibrous terminal bronchioles and forming tubercles there, in addition to the usual foci beneath the pleura. Such tubercles healed in a few months.

#### SUSCEPTIBILITY TO NON-TUBERCULOUS INFECTIONS

There was no specific experiment concerning the effect of inhaled asbestos dust on nontuberculous infection. Intercurrent pneumonia was rather common among animals exposed to asbestos dust, the frequency in guinea pigs exposed in the four inhalation experiments ranging from 16 to 39 per cent. This incidental evidence suggests the possibility of an effect of asbestos dust on nontuberculous infection. Nevertheless, since such epidemics are not uncommon in inhalation experiments with other dusts and even in the colony of normal animals, it is felt that the inhalation of asbestos dust does not exert a significant effect on the susceptibility to nontuberculous pulmonary infection.

#### COMMENT AND SUMMARY

Owing to the vast amount of data included in this investigation, it seems most convenient to summarize and to state as concisely as possible the various observations which emerged from the experiments and to follow each with a brief résumé of the evidence.

A. Various species of animals, including the guinea pig, the rat and the rabbit, but not the mouse and the dog, develop peribronchovascular fibrosis of the lung similar to human asbestosis after being exposed by inhalation or intratracheal injection to long chrysotile asbestos fibers.

Both inhalation and injection experiments provide ample support for this statement. Figure 8 d reveals the cellular fibrosis that occurs in guinea pigs following inhalation of long fiber asbestos; figure 9 shows the fibrosis caused in the rat by inhalation of long fiber asbestos dust. Similar but less extensive fibrosis occurred also in rats and rabbits (table 1). Mice and dogs failed to respond. This variation in response of different species to identical dust exposures is still to be accounted for. B. Long asbestos fibers are essential in the production of the peribronchovascular fibrosis; short fibers are incapable of producing this reaction.

Inhalation experiments with asbestos dust suggest, and intratracheal injection experiments confirm, that peritoneal fibrinosis is produced by asbestos fibers between 20 and 50 microns in length but not by particles shorter than 20 microns (Tables 16 and 18). This indicates that the minimum length of fiber possessing the capacity to produce the typical peritoneal fibrinosis in animals is somewhere between 20 and 50 microns. Panted studies have not been carried out to determine the upper limit of effective fiber length. It appears, however, that that limit will be determined by the inhalability of the fiber.

C. The mode of action of the long asbestos fiber in the production of asbestosis is primarily mechanical rather than chemical in nature.

The evidence for this conclusion has been reviewed in a preceding section, page 39. The flexible filamentary structure of asbestos fibers plays an essential part in the irritating action, since the solid, inflexible filaments of glass wool do not produce fibrinosis (fig. 13 B).

D. Typical experimental asbestosis was produced by the inhalation of an atmospheric suspension containing an average of 138 million asbestos particles per cubic foot of air by light field count, of which less than 1 per cent consisted of fibers longer than 10 microns.

In the inhalation experiment with 100 per cent ball-milled asbestos dust containing 0.6 per cent of fibers longer than 10 microns (table 11) typical fibrinosis was obtained (table 9). The evidence presented shows at least that an atmospheric concentration of asbestos dust containing less than 1 million (0.6 per cent  $\times$  138 million) fibers longer than 10 microns per cubic foot of air is capable of producing experimental asbestosis in guinea pigs. The actual lower limit of concentration of long fibers necessary to produce asbestosis in animals cannot be established from these studies.

E. The duration of exposure required to develop the pulmonary reaction to inhaled asbestos dust is inversely proportional to the concentration of long fibers in the atmosphere; as the concentration is increased, the reaction develops in shorter time.

The basis for this statement appears in the data of the inhalation experiment with long fiber asbestos. For that experiment the average concentration of the atmospheric dust was about 40 million particles per cubic foot of air, and size-frequency determinations disclosed that 6.7 per cent of the air-suspended material consisted of fibers longer than 10 microns (table 11). Thus, by calculation, it is estimated that the concentration of the longer fibers was 2.7 million (6.7 per cent  $\times$  40 million). The lungs of animals exposed to the long fiber asbestos dust revealed that the pulmonary reaction developed in approximately one-half the exposure time required for its development in animals inhaling the ball-milled product, for which the concentration of the longer fibers was only 0.8 million (0.6 per cent  $\times$  138 million).

F. Established experimental asbestosis ceases to progress on discontinuance of dust exposure.

The experimental investigation shows, in fact, that on discontinuance of exposure there was an appreciable clearing of the mature pulmonary

lesions, due to contraction of the fibrous tissue. In contrast, an immature tissue response, evidenced primarily by cells with little or no fibrosis, continued to progress. It is assumed that, following attainment of fibrotic maturity, the same process of contraction would ensue as was noted for the mature lesion.

G. The formation of asbestosis bodies represents a coating of the fibers by blood and tissue elements, which results in loss of ability of the fiber to produce fibrinosis.

Intratracheal injection of asbestosis bodies failed to produce the typical asbestotic tissue reaction in experimental animals. The cessation of progressive reaction observed soon after exposure terminates may be due to the formation of asbestosis bodies.

H. Aluminum hydroxide failed to neutralize the fibrogenic action of the long fiber asbestos.

Aluminum hydroxide added to the suspension of chrysotile asbestos prior to intratracheal injection did not retard or prevent the development of asbestosis in rats.

I. Inhalation of asbestos dust did not alter significantly the final outcome of experimental tuberculosis in two series of guinea pigs exposed to the dust.

The apparently mild influence of asbestos dust is in distinct contrast to the stimulating effect exerted by inhaled quartz on a tuberculous process in the lung. The interpretation must remain tentative, however, since it is based on an investigation limited to two series of guinea pigs exposed to only one kind of asbestos, namely, King's flints. Table 4 shows that when the infection was coincidental with the onset of dust exposure, there was temporary progression of the infectious process, with subsequent healing; when infection was initiated after 26 months of dust exposure, the course of the tuberculosis was not appreciably altered. The latter finding is quite different from our usual experience with quartz dust or with mixed dusts containing quartz, wherein the adverse influence of quartz on a tuberculous infection is manifested most strikingly when infection is initiated after a period of dust exposure, viz., superimposed on a background of established silicosis. As indicated above, this more sensitive test, when applied to asbestos dust, failed to demonstrate that the latter had an adverse influence on a tuberculous infection. The inability of asbestos dust in that experiment to affect unfavorably the tuberculous process furnishes strong support for the interpretation that inhaled asbestos dust has no more than a mildly unfavorable effect on pulmonary tuberculosis.

Acknowledgment is gratefully made to the group of companies of the asbestos industry whose generous financial support made this investigation possible.



# KEASBEY & MATTISON COMPANY

AIR MAIL

MANUFACTURERS OF ASBESTOS AND MAGNESIA PRODUCTS SINCE 1873

AMBLER, PENNSYLVANIA

May 11, 1951

K&M No. 6116

W. W. F. Shepherd, Esq.  
Turner & Newall Ltd.  
77-79 Fountain Street  
Manchester

Dear Mr. Shepherd:

Re: Asbestosis Report

Attached please find copy of my letters of May 10th to both Mr. Roland Starkey and Mr. Charles H. Jackson, as well as copy of the Saranac Laboratory's report entitled "Experimental Studies of Asbestosis."

During the course of my visit to Manchester, between April 16th and 20th, I handed two copies of the printed report to Mr. Hanson who advised that T.A.C.Co. Ltd. would retain and study one copy and that he would pass the other copy along to the Washington Chemical Co. I also handed a copy to Mr. R. G. Scothill for T.B.A.Co. Ltd.

Should you or anyone else in the T&N organization require additional copies, please let me know, because we have a supply on hand.

Yours sincerely,

Keasbey & Mattison Company

em:ck  
encls.

cc By Boat

COPY LETTER FROM  
KEASBEY & MATTISON COMPANY  
AMBLER, PA.

May 10, 1951

K&M No. 23 - AAC

Mr. Chas. H. Jackson  
Atlas Asbestos Company Ltd.  
5600 Hochelaga Street  
Montreal

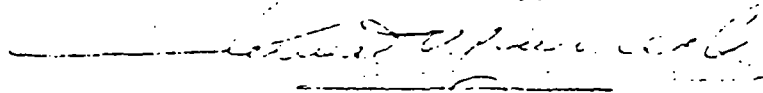
Dear Mr. Jackson:

Some years ago the Asbestos Industry in the United States contracted with the Saranac Laboratories at Saranac Lake, New York, to carry out experiments and issue a report on asbestosis.

This report has finally been issued and I have pleasure in handing you herewith one copy of the reprint from the American Medical Association, Archives of Industrial Hygiene and Occupational Medicine, January 1951.

With kind regards, I remain

Yours sincerely,



Keasbey & Mattison Company

em:ck  
encl.

cc WWFS

May 10, 1951

Roland Starkey, Esq.  
Rhodesian & General Asbestos Corp.  
P. O. Box 1100  
Bulawayo, So. Rhodesia

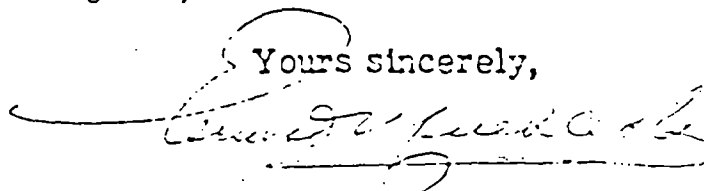
Dear Mr. Starkey:

Some years ago the Asbestos Industry in the United States contracted with the Saranac Laboratories at Saranac Lake, New York, to carry out experiments and issue a report on asbestosis.

This report has finally been issued and I have pleasure in handing you herewith one copy of the reprint from the American Medical Association, Archives of Industrial Hygiene and Occupational Medicine, January 1951.

With kind regards, I remain

Yours sincerely,



Keasbey & Mattison Company

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