Non-malignant asbestos pleural disease

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ABSTRACT During a 10-year period (1970-79) all patients in Uppsala County found to have pleural changes related to asbestos exposure were followed. The lesions could be divided into four types: parietal pleural plaques, exudative pleurisy, thickening of the visceral pleura, and progressive pleural fibrosis. There were 891 cases. The most common type was parietal plaques, which was seen in 827 patients, some of whom later developed other changes. In 22 patients exudation was proven radiologically, and in 84 more cases obliteration of the costo-phrenic angle was seen. The exudations almost all had a benign course, despite sometimes fairly large and bloody effusions. They were practically all symptom-free, being a surprise finding on chest radiography. Thickening of the visceral pleura can only be seen radiologically in the fissures and occurred in a few cases in addition to other changes. In a small group of more heavily exposed individuals, a progressive pleural fibrosis developed, sometimes after an initial effusion.

Pleural changes are common among those exposed to asbestos. Most common are plaques on the parietal pleura, which can produce radiographic changes virtually pathognomonic of exposure to asbestos. Unspecific abnormalities, such as exudation and pleural thickening, can also occur. Diagnosis of these changes depends upon a history of asbestos exposure and exclusion of other causes.\(^1\) A study of asbestos-related pleural changes has been in progress in the county of Uppsala for a 10-year period (1970-79).

Methods

The county of Uppsala is situated just north of Stockholm. There are approximately 250,000 inhabitants, almost half of whom live in Uppsala, a University town with comparatively few industries. There are no dock yards, asbestos factories, or similar former heavy users of asbestos in the county. Persons with asbestos exposure are mainly builders, plumbers, and so on or have worked in small scale manufacturing shops.

A general health survey has been in practice in the county since 1960. This includes a 100 × 100 mm size chest radiograph. All inhabitants from age 15 years and above are invited to participate every second or third year, and the average attendance is 70%.\(^2\) Thus, for many of the inhabitants serial films over a 20-year period are available.

Since 1970, all residents in the county found to have pleural changes thought to be caused by asbestos have been followed. Most have been found in the health survey, but some cases (less than 5%) have also been referred to the clinic.

The pleural changes were divided into four types on the basis of radiology (table 1), but two or more types often occurred together in the same patient. The diagnostic criteria were as follows: Pleural plaques were diagnosed only if they were bilateral and of at least grade II according to ILO/UICC—that is, at least 5 mm thick when seen tangentially.\(^3\) Asbestos exposure was not a prerequisite, since exposure can be very difficult to ascertain. Pleural effusions alone were not included unless plaques were present before or developed after the effusion appeared. For such cases significant asbestos exposure was required, and other causes of the effusion had to have been excluded. Only clinically motivated investigations were performed. Thickening of the visceral pleura can only be seen radiologically in the fissures and was diagnosed only in connection with other changes. Progressive pleural fibrosis was thought to be caused by asbestos if there was a history of such exposure and other causes had been excluded. Exposure to asbestos was graded from 1 to 3 on the basis of the patient's history. Grade 1 was intermittent, not very heavy exposure, grade 3 heavy daily use, and grade 2 in between.
Table 1  Non-malignant asbestos-related pleural changes: classification

<table>
<thead>
<tr>
<th>Afflicted pleura</th>
<th>Appearance</th>
<th>Development</th>
<th>Typical X-ray</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parietal pleural plaques</td>
<td>Parietal</td>
<td>Very gradual</td>
<td>Clear angles, apices not affected</td>
</tr>
<tr>
<td>Uncomplicated pleural exudate</td>
<td>Both</td>
<td>Sudden</td>
<td></td>
</tr>
<tr>
<td>Thickening of visceral pleura</td>
<td>Visceral</td>
<td>Very gradual</td>
<td>Seen only in fissures, mainly in minor fissure</td>
</tr>
<tr>
<td>Progressive pleural fibrosis</td>
<td>Both</td>
<td>Can start suddenly</td>
<td>Sinuses always involved &quot;Crow's feet&quot; common early sign, Apices sometimes affected</td>
</tr>
</tbody>
</table>

Results

There was a total of 891 persons with pleural changes associated with exposure to asbestos.

Parietal pleural plaques

These were by far the most common, 827 patients showing such changes some time during the observation period, some of them later developing other types as well. The incidence of new cases was growing each year, and it could be estimated that at the end of 1979, at least 2.2% of the men 40 years of age or older had radiological pleural plaques. The plaques were very rare in women and below age 40 years.

More than 90% were asked about exposure to asbestos, and of those, 80% confirmed exposure. In another 10%, exposure was probable from the occupation. The average time from first exposure to first radiograph fulfilling the criteria was 30 years. The plaques showed a slow and gradual development and could often be suspected on earlier films. They have been thoroughly described in an earlier paper.

The "average grade of exposure" for pleural plaques only was 1.4, if those denying exposure were excluded.

Uncomplicated pleural effusion (figs 1, 2)

Effusion, proven radiologically and in most cases also by aspiration, was found in 22 patients. In four of these the effusion developed into progressive pleural fibrosis (table 2). The exudate tended to persist for some months or up to a year, and sometimes recurred on the same or the other side after one or two years. The fluid was often macroscopically bloodstained and microscopically there was a predominance of erythrocytes or mature lymphocytes or both. Up to three aspirations were performed with 50 to 1500 ml removed (table 3).

As seen in table 2, pleural plaques were present before the effusion or developed afterwards. It is

Fig 1  a: 1972. Right-sided effusion, haemorrhagic at aspiration. No plaques seen. b: 1979, same patient. Rounded angle right, typical plaques left.
probable that in some cases plaques developed later but were obscured by the remnants of the pleurisy. The plaques seemed in no way affected by the effusion, except for being hidden in the fluid and reappearing when it was resolved or in some cases remaining obscured by pleural thickening afterwards.

Sedimentation rate was usually normal but in a few cases raised to up to 50 mm. Slight leukocytosis was seen only occasionally. There were no signs of collagen diseases. Cytology of the fluid and biopsy of the pleura sometimes gave reason to suspect a tumour but all patients have been followed for at least three years with no signs of neoplasm (table 3).

Table 2 Pleural effusion in 22 cases

<table>
<thead>
<tr>
<th>Course</th>
<th>Number of cases</th>
<th>Special notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Disappeared without trace; years later pleural plaques</td>
<td>4</td>
<td>Persisted 2-5 months.</td>
</tr>
<tr>
<td>Effusion in persons with pleural plaques; disappeared without trace</td>
<td>10</td>
<td>Recurrence same side, 2 years later in one case, Persisted up to 6 months. Two recurrences 2 years later—one same side, one opposite. Persisted 2 and 4 months respectively.</td>
</tr>
<tr>
<td>Left rounded angle; years later pleural plaques on opposite side</td>
<td>2</td>
<td>Persisted 4 months</td>
</tr>
<tr>
<td>Effusion in persons with pleural plaques, leaving rounded sinus</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Developed into progressive pleural fibrosis</td>
<td>4</td>
<td></td>
</tr>
</tbody>
</table>

Typically, there were no or very slight symptoms. Only one patient had a chest film because of symptoms (general malaise and slight pain in the side). All the others were found accidentally and admitted only vague symptoms on questioning. Average grade of exposure was 1-3, and the mean “latency time” around 30 years, with a wide variation of nine to 46 years.

Obliteration of a costophrenic angle with or without some degree of pleural thickening without a known episode of pleural exudate was observed in 84 cases. In 28 cases, it occurred before development of pleural plaques. Depending on the degree of residual pleural thickening, the plaques were seen on the contralateral side only (15 cases) or on both sides (13 cases). In 31 cases sudden obliteration of the angle was noticed on follow-up of pleural plaques. In only five cases was the obliteration accompanied by a considerable thickening of the pleura, but in all five this was still unchanged some years later. Twenty-five patients had bilateral costa-phrenic angle obliteration.

Average “latency time” was 34 years. Very few patients had noticed any symptoms since the earlier film with a normal angle. The average grade of exposure to asbestos in patients with one or both angles obliterated was 1-4.

Thickening of the visceral pleura (fig 3)
This was not very common and tended to occur with comparatively heavy exposure, but was not systematically looked for until the last few years.
Table 3  Benign asbestos pleurisy. Effusions of at least 400 ml at thoracocentesis. Some clinical data

<table>
<thead>
<tr>
<th>Side</th>
<th>Pre-effusion X-ray</th>
<th>Years from first exposure</th>
<th>Grade of asbestos exposure*</th>
<th>Total amount tapped (ml)</th>
<th>Times tapped</th>
<th>Macroscopic appearance</th>
<th>Remnants on X-ray</th>
<th>Observation since exudate (years)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R</td>
<td>Plaques</td>
<td>45</td>
<td>1</td>
<td>450</td>
<td>1</td>
<td>Bloody</td>
<td>No trace</td>
<td>3</td>
</tr>
<tr>
<td>R</td>
<td>&quot;</td>
<td>51</td>
<td>2</td>
<td>500</td>
<td>1</td>
<td>&quot;</td>
<td>&quot;</td>
<td>3</td>
</tr>
<tr>
<td>L</td>
<td>&quot;</td>
<td>22</td>
<td>1</td>
<td>500</td>
<td>2</td>
<td>&quot;</td>
<td>&quot;</td>
<td>5</td>
</tr>
<tr>
<td>L</td>
<td>Normal</td>
<td>50</td>
<td>1</td>
<td>600</td>
<td>2</td>
<td>&quot;</td>
<td>Bilateral progressive fibrosis</td>
<td>5</td>
</tr>
<tr>
<td>R</td>
<td>&quot;</td>
<td>51</td>
<td>1</td>
<td>300</td>
<td>1</td>
<td>&quot;</td>
<td>&quot;</td>
<td>Same patient</td>
</tr>
<tr>
<td>L</td>
<td>Plaques</td>
<td>18</td>
<td>1</td>
<td>850</td>
<td>3</td>
<td>&quot;</td>
<td>No trace</td>
<td>6</td>
</tr>
<tr>
<td>R</td>
<td>Normal</td>
<td>30</td>
<td>2</td>
<td>550</td>
<td>1</td>
<td>&quot;</td>
<td>Obliterated sinus Plaques other side 3 years later Obliterated sinus</td>
<td>7</td>
</tr>
<tr>
<td>R</td>
<td>Plaques</td>
<td>17</td>
<td>1</td>
<td>1000</td>
<td>1</td>
<td>&quot;</td>
<td>No trace, three years later plaques</td>
<td>8</td>
</tr>
<tr>
<td>R</td>
<td>Normal</td>
<td>33</td>
<td>3</td>
<td>2000</td>
<td>3</td>
<td>Yellowish</td>
<td>&quot;</td>
<td>18</td>
</tr>
</tbody>
</table>

*See text.

Progressive pleural thickening (fig 3)

This occurred in 27 patients. The costophrenic angles were involved in all. In four it started with an effusion, as mentioned above. The typical finding was progression when observed over a few years. The speed of progression varied. The lesion was always bilateral, even if it often started unilaterally. The first signs were often streaks radiating towards or away from the periphery (“Crow’s feet” as described by Mackenzie). These probably represent local scarring or convolution of the pleura, and they were also sometimes seen as a non-progressive remnant of pleural effusion. As the disease advanced, these changes became confluent. Unlike all other types, considerable thickening of the pleural cap was sometimes noted.

For various reasons, pleurectomy was performed on one or both sides of four patients. In only one of them was there subjective improvement in dyspnoea. This lack of improvement was caused by coexistent pulmonary fibrosis, proven by pulmonary biopsy.

Sixteen patients started with pleural plaques as the sole abnormality, then developed first unilateral and later bilateral obliteration of the sinuses, and finally extensive thickening of the pleura.

In four cases, calcified plaques were observed through the pleural fibrosis. In three of these, the calcification occurred after the hyaline plaques had been obscured by the fibrosis, indicating that the plaques continue their development independently of other pleural changes.

The latency time was 34 years, and the average exposure to asbestos was 2-0, higher than in the other groups.

Pseudotumour formation

In six patients, three after effusion and three with progressive thickening, a “pseudotumour” developed—that is, a lesion which on radiography was strongly suggestive of tumour but at operation proved to be visceral pleural thickening and “folded lung”.

Discussion

Pleural changes are common in pulmonary asbestosis. In advanced cases they are mainly progressive thickening, as described here. The explosive increase in the use of asbestos during and after world war 2 led to a large number of persons being exposed in fairly low doses. With a latency time of three decades, the result of this is now seen and will be seen increasingly in the near future.

At least 2.2% of men in Uppsala County above age 40 years have radiologically visible pleural plaques and many more at necropsy and the figure is rising. In areas with dockyards, asbestos factories, and so on the figure will be much higher, and other pleural changes caused by asbestos should also be more common than with the relatively low grade of exposure in Uppsala County.

Pleural effusion is a well-known sequel to asbestos exposure (table 4). It has been stated that the underlying lung is invariably abnormal, but in the present series most cases run an uncomplicated course without signs of lung disease. Judging from table 4, mesotheliomas are a common late complication, or the exudate might be the first sign of the tumour. Three years of observation, as in the first two cases of table 3, is a short period to exclude a mesothelioma, but the absence of symptoms or radiological signs is against this diagnosis. The cases in table 4 are probably heavily selected explaining the high incidence of tumour.

Pleural effusion caused by asbestos exposure is
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often asymptomatic. Of the 22 proven effusions in this series, all but one were discovered accidentally. Only 10 of them—that is, less than half—left radiologically visible remnants. Thus, those exudates that are diagnosed are only the top of the iceberg.

Apart from tuberculosis, collagen diseases, and drug-induced reactions, mesothelioma is the main differential diagnosis. Vigorous investigation to prove or disprove this diagnosis seems unwarranted in view of the disappointing results of treatment of this tumour.

Thickening of visceral pleura occurs in many patients exposed to asbestos and is easily demonstrated by the pathologist but difficult for the radiologist to see, except in the fissures. The pleura gradually thickens with time and finally calcifies. Calcification can also be seen in the major fissures (Sargent EN, personal communication 1980). Thus, there is a striking parallel with the plaques in the parietal pleura, the main difference being that the visceral thickening is more diffuse and therefore more difficult to see on radiographs.

Fig 3  a: 1969. Rounded left sinus, pleural plaque laterally on right. b: Same patient, 1973. Left, extensive pleural changes. Right, plaque has grown and interlobar fissure has thickened (type 3). c: Same patient, 1978. Pleurectomy on left side; on the right, rounded angle and pleural fibrosis with crows’ feet.
Table 4  Benign asbestos exudates: literature review

<table>
<thead>
<tr>
<th>Author</th>
<th>Number of cases</th>
<th>Latency mean (years)</th>
<th>Time (years) limits</th>
<th>Symptoms</th>
<th>Complications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Boutin et al(^5)</td>
<td>3</td>
<td>44</td>
<td>42-49</td>
<td>One pleuritic pain</td>
<td></td>
</tr>
<tr>
<td>Christen et al(^6)</td>
<td>10</td>
<td>26</td>
<td>10-38</td>
<td>Insidious</td>
<td>Pleuritic pain</td>
</tr>
<tr>
<td>Eisenstadt(^7)</td>
<td>1</td>
<td>—</td>
<td>—</td>
<td>Pleuritic pain</td>
<td></td>
</tr>
<tr>
<td>Eisenstadt(^8)</td>
<td>3</td>
<td>—</td>
<td>—</td>
<td>Two pain, one also fever</td>
<td></td>
</tr>
<tr>
<td>Elder(^9)</td>
<td>13</td>
<td>—</td>
<td>—</td>
<td>Two mesotheliomas (after 10</td>
<td>One mesothelioma after 12 years</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>and 14 yrs respectively)</td>
<td></td>
</tr>
<tr>
<td>Gaensler and Kaplan(^10)</td>
<td>12</td>
<td>20</td>
<td>4-46</td>
<td>Four pain, 9 dyspnoea</td>
<td></td>
</tr>
<tr>
<td>Léménager et al(^11)</td>
<td>5</td>
<td>22</td>
<td>7-46</td>
<td>Two pain and dyspnoea</td>
<td></td>
</tr>
<tr>
<td>Mattson(^12)</td>
<td>11</td>
<td>18</td>
<td>6-29</td>
<td>Four slight pain</td>
<td></td>
</tr>
<tr>
<td>Navratil and Dobias(^13)</td>
<td>7</td>
<td>—</td>
<td>—</td>
<td>Pain in some cases</td>
<td></td>
</tr>
<tr>
<td>Nyiredy(^14)</td>
<td>1</td>
<td>14</td>
<td>—</td>
<td>Pain</td>
<td></td>
</tr>
<tr>
<td>Sluis-Cremer and Webster(^15)</td>
<td>9</td>
<td>14</td>
<td>6-36</td>
<td>Three pain, 2 dyspnoea</td>
<td></td>
</tr>
<tr>
<td>Smyth(^16)</td>
<td>1</td>
<td>45</td>
<td>—</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>75</td>
<td>22</td>
<td>4-49</td>
<td>Approximately 20 pain, 10</td>
<td>Nine mesotheliomas, 2 cancers,</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>dyspnoea</td>
<td>4 deaths of “pleural hyalnosis”</td>
</tr>
</tbody>
</table>

Progressive pleural fibrosis has been described by many authors.\(^5\)\(^7\)\(^14\)\(^25\) Wright et al\(^26\) believed the patients’ dyspnoea to be caused by restrictive pleural fibrosis compressing the lung, but fibrosis of the lung itself probably often coexists, making pleurectomy of doubtful value. Important differential diagnoses are drug reactions and mesothelioma. The prognosis of these lesions is variable. In some patients we now have a follow-up of a decade or more with only slight handicap and slow progression, while others rapidly develop severe dyspnoea.

“Atelectatic pseudotumours” may be caused by fibrotic changes in the pleura of any cause, leading to deformation of the underlying lung and a bronchographic picture reminiscent of “trees bent by the wind”.\(^27\)\(^28\) In asbestosis this has been described as “broom deformation”.\(^16\)\(^28\) If this deformation causes kinking of some bronchi, a “rounded atelectasis” can occur. The lung can also become atelectatic while floating in an effusion and by adhesions remain so when the exudate is absorbed. These pseudotumours are not uncommon in persons exposed to asbestos.\(^30\)

References

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