Decline in annual lung function in workers exposed to asbestos with and without pre-existing fibrotic changes on chest radiography

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Abstract

Objectives—To examine whether or not workers with pre-existing mild pulmonary fibrosis have accelerated decline in forced expiratory volume in one second (FEV₁) or forced vital capacity (FVC), under low level exposure to chrysotile asbestos.

Methods—All male workers in two asbestos manufacturing factories were followed up annually for six years to compare their declines in FEV₁ and FVC. The values of FEV₁ and FVC were divided by the square of the person’s height to adjust for body size differences (FEV₁/Hᵗ² and FVC/Hᵗ², respectively). Annual change was calculated for each subject as a slope of the simple linear regression with FEV₁/Hᵗ² or FVC/Hᵗ² regressed according to age. Analysis was conducted on 242 middle aged workers who had normal routine spirometry values, normal chest radiographs or mild pneumoconiosis up to 1/2 grade, without changes either in smoking habit or severity of pneumoconiosis during the study period, and with acceptable spiromograms in three or more surveys. The occupational environment, in terms of chrysotile asbestos, had been well controlled below the threshold limit value of Japan at that time—namely, 2 fibres/ml.

Results—There was no significant effect from the interaction between pre-existing mild pulmonary fibrosis and a low level of exposure to chrysotile asbestos on the accelerated annual decline of FEV₁/Hᵗ² or FVC/Hᵗ². Fibrosis significantly contributed to annual changes in FEV₁/Hᵗ², even after adjustment for mean FEV₁ and smoking. The point estimate of the contribution was -4.9 ml/m³/y. No significant independent contribution of exposure was found in decline of either FEV₁/Hᵗ² or FVC/Hᵗ².

Conclusions—Pre-existing pulmonary fibrosis is an independent risk factor for accelerated annual decline of FEV₁, even when mild and stable. Additional decline due to exposure to chrysotile asbestos is less probable if it is well controlled under the current threshold limit value.

Keywords: pneumoconiosis; asbestos; spirometry

Recent advances in industrial hygiene techniques have remarkably improved the occupational environment, and currently actual exposure to asbestos and other mineral dusts and fibres is generally controlled at a low level. The number of new cases with severe fibrotic changes in the lungs has been decreasing in Japan and most other developed countries.

In contrast, considerable concern has come to be directed towards work related chronic obstructive pulmonary disease (COPD), one of the important outcomes of inhaling harmful substances under such low exposure conditions. Accelerated annual loss of forced expiratory volume in one second (FEV₁) has been reported in several longitudinal studies on coal miners, gold miners, and asbestos cement workers. Little is known about the effect of pre-existing pulmonary fibrosis on the subsequent annual decline of FEV₁ under the current low exposures to mineral dusts and fibres.

The purpose of this analysis was to examine the synergistic effect of low exposure to chrysotile asbestos and pre-existing mild pulmonary fibrosis, on the mean annual decline in FEV₁ and forced vital capacity (FVC), together with their independent effects on these two indices.

Subjects and methods

Subjects
Study subjects were recruited from two asbestos manufacturing factories where calcium silicate boards and joint materials including asbestos were produced. All male employees of the factory were first examined in 1985 by spirometry, chest radiography, and questionnaire. They were followed up annually up to 1991 (excluding 1987) with the same protocol as used in the first assessment. Of this cohort, we limited our scope of analysis to those who had worked in these factories for three years or more between 1985 and 1991 to obtain a reliable estimate of annual change in spirometric indices. Thus the original sample was 351 subjects. The age range of the cohort at the beginning of this study was from 19 to 58.

Spirometry
The procedure used has been described in detail elsewhere. Briefly, spiromograms were measured with a dry rolling seal spirometer (Chestac 65, Chest Co, Japan). Subjects, in a standing position, were asked to repeat the forced expiration manoeuvre at most seven
times to obtain acceptable, reproducible results. Nose clips were not used because an open circuit method was adopted. Usual body temperature, pressure, and saturation correction and back extrapolation correction were carried out. Whether or not each manoeuvre was acceptable was evaluated by the following criteria: starting without hesitation, apparent maximal effort, and smooth continuous exhalation without cough. Reproducibility was judged by the criteria of the American Thoracic Society,7 based on the measured values, and on the shapes of flow-volume and flow-time curves. In each survey, the subjects with at least two reproducible spirograms were considered acceptable. The figures of FEV1 and FVC used in this analysis were derived from the best manoeuvre with the largest sum of FVC and FEV1. To minimise the measurement bias across the surveys, all examinations were carried out by one examiner with the same spirometer throughout all the surveys.

CROSS SECTIONAL AND LONGITUDINAL INDICES
At first, we examined several models for FEV1 and FVC to different powers of height, to standardise the value according to body size. The results showed that the height squared (Ht2) model seemed to be appropriate in terms of this standardisation. We then used the Ht2 proportional values of FEV1 and FVC (FEV1/Ht2 and FVC/Ht2, respectively) in this analysis, to adjust for height differences. Annual changes in FEV1/Ht2 and FVC/Ht2, a longitudinal index denoted by “slope” in this paper, were calculated for each subject as the slopes of simple linear regressions in which FEV1/Ht2 and FVC/Ht2 were regressed by age. A cross sectional index, FEV1/Ht2 level—the mean FEV1/Ht2 value of each subject during the follow up period—was used. The first FEV1/Ht2 value for each worker was also considered, but it seemed inappropriate because of the relatively large error in FEV1, each time it was measured in comparison with the size of the decline in FEV1. By using mean values, the error of each measurement could be averaged.

OCCUPATIONAL EXPOSURE
Among primary sources of exposure to asbestos during the follow up were exposure to the processing of chrysotile asbestos in the production of calcium silicate boards, some joint and sealing materials, and insulating products. Periodic monitoring of the occupational environment has been carried out not only in all work areas where asbestos was used, but other possible areas such as storage and stock yards. Based on records made during the study period, the time weighted average (TWA) concentration of chrysotile asbestos fibres has been controlled to below 0.5 fibres/ml, well below the permissible levels of occupational exposure in Japan—namely, two fibres/ml for chrysotile.8 The company records suggested that higher exposure might have been probable in the past, at least before the mid-1970s. For this paper, however, I could not obtain either a reliable estimate of cumulative exposure in the past or a personal exposure level of each worker during the study period. Therefore, according to the actual work done during the follow up period, subjects were divided into two groups. Those who had worked for more than one year in work areas where asbestos was processed, were considered to be exposed, and others were unexposed.

FINDINGS OF PNEUMOCONIOSIS
Chest radiography was carried out with standard procedures for Japan.9 Severity of pneumoconiosis in lung fields was classified into 12 grades from 0/− to 3/+ . According to the findings before the first survey and during the follow up period, subjects who continued to be classified as 0/− , 0/0, or 0/1 were considered negative, and those who had been classified as 1/0, 1/1, or 1/2 were considered positive. Those who had been classified as 2/1, or more, were excluded from analysis. Furthermore, any subject who had moved from negative to positive or had shown a deterioration in the severity of pneumoconiosis during follow up, was also excluded from analysis. Pleural findings were not considered in this analysis.

QUESTIONNAIRE SURVEY
The standardised questionnaire of the American Thoracic Society (ATS-DLD-78-A)10 was used with slight modification and translation into Japanese to assess smoking habit, as well as respiratory symptoms and medical histories. Mean cigarette consumption during the follow up period was calculated for each current smoker as the number of cigarettes smoked a day. For non-smokers, this was defined as zero.

ANALYSIS
All statistical tests and estimations were carried out with the SAS statistical packages at the Tokyo University Computer Center. Interaction of pneumoconiosis and exposure, and independent effects of each of these factors were estimated with an analysis of covariance (ANCOVA) in which the mean level of FEV1/Ht2 and smoking were treated as covariates.

Results
In 323 subjects out of the original 351 (92%) forced spirometry was carried out acceptably three times or more and the effective duration of follow up was three years or more. Of these, we excluded 18 subjects who had changed their smoking habit during the follow up and nine whose chest x ray film classification had been considered to be severe or who deteriorated during the study. There were 17 subjects with lowered routine spirometric results (the percentage predicted value of vital capacity calculated by the prediction equation used in Japanese pneumoconiosis law): VC (%) < 80 or the ratio of FEV1 (%) to FVC < 70 at their first examination. According to
the pneumoconiosis law in Japan, workers exposed to dust with reduced pulmonary function (as already described) are recommended to be transferred to work free from exposure to dust. Therefore the inclusion of subjects with reduced VC (%) or FEV₁ (%) in the analysis may have caused a selection bias for the objectives of this study. Also, they could have shown variable aging patterns in pulmonary function due to this deterioration. For these reasons workers with low spirometric results were excluded from the analysis. I also excluded 38 subjects aged less than 30 at the first examination because their age related changes in FEV₁ and FVC might have been different from those of the other middle aged subjects as a steady decline in FEV₁/Ht² seemed to occur after 30 years of age. Actually the mean (SD) FEV₁/Ht² was −0.0 (10.3) ml/m² in the 19–29 year old group, and −9.8 (9.8), −8.3 (11.4), and −10.0 (12.5) in the 30–39 year old, 40–49 year old, and 50 year and over groups, respectively. So we analysed 242 middle aged subjects who had not shown changes in either their smoking habit or severity of pneumoconiosis during the follow up period, who had normal routine spirometric results at the first examination. Table 1 shows the mean (SD) of the slopes of FEV₁/Ht² and FVC/Ht², and compares those included in the analysis and those excluded.

Table 1: Crude comparison of mean (SD) annual decline of FEV₁, and FVC (adjusted for height squared (Ht²)) between those included in and excluded from analyses

<table>
<thead>
<tr>
<th></th>
<th>FEV₁/Ht² (ml/m²/y)</th>
<th>FVC/Ht² (ml/m²/y)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Those analysed</td>
<td>−9.1 (11.2)</td>
<td>−5.5 (13.7)</td>
</tr>
<tr>
<td>Those excluded</td>
<td>81*</td>
<td>−6.0 (10.2)</td>
</tr>
<tr>
<td>&lt;30 years old</td>
<td>38</td>
<td>−13.5 (10.8)</td>
</tr>
<tr>
<td>Change in smoking habit</td>
<td>18</td>
<td>−10.2 (12.1)</td>
</tr>
<tr>
<td>Low spirometry</td>
<td>17</td>
<td>−13.8 (15.6)</td>
</tr>
<tr>
<td>Change in chest radiographs</td>
<td>9</td>
<td>−10.2 (16.7)</td>
</tr>
<tr>
<td>Total acceptable respondents</td>
<td>323</td>
<td>−7.4 (15.4)</td>
</tr>
</tbody>
</table>

*One person had two reasons for being excluded.

Table 2: Mean (SD) of related variables

<table>
<thead>
<tr>
<th>Pneumoconiosis</th>
<th>Negative exposure to asbestos</th>
<th>Positive exposure to asbestos</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(−) (n = 90)</td>
<td>(+) (n = 109)</td>
</tr>
<tr>
<td>Age (y)</td>
<td>43.3 (5.4)</td>
<td>43.6 (6.2)</td>
</tr>
<tr>
<td>Smoking:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Smokers (%)</td>
<td>56.7</td>
<td>69.4</td>
</tr>
<tr>
<td>Cigarettes/day (n)</td>
<td>20.0 (7.5)</td>
<td>18.4 (5.0)</td>
</tr>
<tr>
<td>FEV₁/Ht² (ml/m²)</td>
<td>1.22 (0.17)</td>
<td>1.20 (0.15)</td>
</tr>
<tr>
<td>FVC/Ht² (ml/m²)</td>
<td>8.55 (5.6)</td>
<td>8.54 (5.6)</td>
</tr>
</tbody>
</table>

Table 3: Simple comparison of means (SD) of slopes of FEV₁/Ht² and FVC/Ht² (ml/m²/y) by exposure and pneumoconiosis

<table>
<thead>
<tr>
<th>Pneumoconiosis</th>
<th>Negative</th>
<th>Positive</th>
</tr>
</thead>
<tbody>
<tr>
<td>FEV₁/Ht² Exposure:</td>
<td>−7.6 (1.1)</td>
<td>−11.3 (3.2)</td>
</tr>
<tr>
<td>PFC/Ht² Exposure:</td>
<td>−3.5 (1.3)</td>
<td>−6.8 (3.1)</td>
</tr>
</tbody>
</table>

*VC was based on the prediction equation used in the pneumoconiosis law of Japan*.

Figure 1: Two way scattergram of the slopes of FEV₁/Ht² in all analysed subjects.
Table 4 shows the effects of age and current smoking on annual changes in FEV/HT and FVC/HT. The effect of age was not clear, but the oldest group showed a slightly faster decline in both of the indices. A dose related acceleration due to smoking was found up to the category of 20 to 29 cigarettes a day.

The relation between the cross sectional level of FEV, and the two longitudinal slopes were examined by regression analysis. Figure 2 shows a significant positive association with FEV/HT level in both change in FEV/HT (P < 0.001) and change in FVC/HT (P < 0.001).

Analysis of covariance was conducted to avoid possible confounding, in which pneumoconiosis and exposure to asbestos were treated as independent factors (fig 3). At first, age, mean cigarette consumption, and FEV/HT level were considered as covariables. Age did not contribute greatly to the ANCOVA model (table 4). So age was then excluded from the analysis. Figure 3 shows the square means (SEMs) of the slopes for pneumoconiosis and exposure to asbestos estimated with this model, adjusted for smoking and level of FEV/HT. Although the least square mean in the exposed group suggested faster annual decline compared with the other three categories, particularly in FVC/HT, there was no significant interactive contribution of the two factors either to the slope of FEV/HT or to that of FVC/HT.

Table 5 shows the independent effects of the factors and covariables estimated simultaneously by ANCOVA models, with the interaction between pneumoconiosis and exposure to asbestos being excluded, based on the results shown in fig 3. Even after adjustment for smoking and FEV/HT level, pneumoconiosis contributed significantly to FVC/HT slope. The point estimate of the effect was -4.9 ml/m²/y for the slope of FEV/HT. This corresponds to an additional annual loss of 13 ml in FEV, for a subject with an average height of 165 cm. A similar tendency was found in the FVC/HT slope, but it was not significant. Mean cigarette consumption, as a
positive control in this analysis, was significantly associated with an accelerated decline of FEV₁, but not with a decline in FVC. There was no significant contribution of exposure to asbestosis on the slopes.

Discussion
The results of recent longitudinal studies in occupational cohorts exposed to mineral dusts and fibres have shown a significant association between such forms of exposure and accelerated annual loss of FEV₁. An important point in this context is that the association could be found even under relatively low exposure conditions. For example, in asbestosis workers Ohlson and coworkers reported accelerated loss of FVC and FEV₁ even in the workers who had retired from asbestos processing under exposures of around one or two fibres/ml. Another important point is the interaction of such low levels of exposure and pre-existing pulmonary fibrosis, because numerous workers who have mild pulmonary fibrosis due to a high level of exposure in the past are still working under the current conditions of low exposure. Therefore, it seems to be important to examine whether or not there is an interaction between current low exposure and pre-existing mild pulmonary fibrosis on deterioration of lung function, together with their independent effects.

In a simple comparison of the slopes of the categories defined by exposure and pneumoconiosis, in this study, the workers with fibrotic findings in the lungs, although their findings were mild and stable, had a tendency to show an accelerated slope compared with those who did not have fibrosis. Furthermore, it seemed that the tendency was more obvious in the exposed subjects. The ANCOVA analysis, however, indicated that the interaction of pneumoconiosis and exposure to asbestosis did not contribute significantly to the slopes, when the two possible confounding factors, smoking and cross sectional level of FEV₁, were taken into consideration. This result suggested that the effects of smoking and FEV₁ level may have acted additively, although the possibility of type 2 error could not be neglected in assessing the interaction due to the few subjects in the exposed group.

The independent effect of pneumoconiosis on decline of FEV₁/Ht² was highly significant, even after adjustment for cigarette consumption and FEV₁/Ht² level. In other words, those workers with pre-existing mild fibrosis had accelerated annual loss in FEV₁/Ht². The independent effect of exposure to asbestos did not reach significance. As the difference in decline of FEV₁, adjusted on cross sectional level indicates the potential causal role of a given factor on decline of FEV₁, pre-existing pulmonary fibrosis would be an independent risk factor for accelerated annual decline of FEV₁, irrespective of exposure, even if it is mild and stable.

One possible source of bias in this analysis would be the selection bias, or the healthy worker effect. It is possible that those susceptible to exposure to asbestos and then inclined to rapid decline in FEV₁, had been transferred from asbestos processing work to other safer or unexposed types of work before this study started. In that case, the exposed workers must have been healthier and less susceptible to exposure than the unexposed group. In other words, at the beginning of the study, the unexposed group might have included more of those with a natural rapid decline in FEV₁ than the exposed group. If this is the case, it might partially explain why no difference in slopes by exposure to asbestos was detected in this study.

Table 2 shows that routine spirometric results of our subjects at the start of this study were good, and the unexposed subjects had rather better results than the exposed at the beginning of the study. Furthermore, our eligibility criteria for lung function would have excluded any subject who's lung function had deteriorated and may have led to work transfer. Also, any worker with fibrosis of grade 2/1 and more was excluded from the analysis. Thus this study included only healthy subjects and subclinical cases and it is therefore unlikely that the healthy worker effect seriously biased our results.

Another important factor is past exposure. We can assume that cumulative exposure accelerates decline in both FEV₁ and FVC independently from the current exposure. Because of this assumption, it is possible that the slope in old workers might be exaggerated. In my study, however, no notable age related difference in slopes was found. Therefore, it seems unlikely that the independent effect of pneumoconiosis has been seriously overestimated, although those workers with pneumoconiosis were slightly older than those without. It is still possible that the presence of fibrosis can simply be a marker of past heavy exposure, not necessarily a marker of susceptibility to accelerated decline in FEV₁. Further analysis will be needed to make this point clear.

Among confounding factors, current cigarette consumption was negatively associated with FEV₁/Ht² slope, which may have acted additively, although the possibility of type 2 error could not be neglected in assessing the interaction due to the few subjects in the exposed group.

Table 5: Independent effects of explanatory variables on annual change of FEV₁ and FVC (adjusted for height squared (Ht²)) estimated simultaneously by analysis of covariance

<table>
<thead>
<tr>
<th>Variable</th>
<th>FEV₁/Ht²</th>
<th>FVC/Ht²</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>SEM</td>
</tr>
<tr>
<td>Pneumoconiosis</td>
<td>-4.9</td>
<td>1.9</td>
</tr>
<tr>
<td>Exposure (pack/day)</td>
<td>0.08</td>
<td>1.4</td>
</tr>
<tr>
<td>Smoking (pack/day)</td>
<td>2.7</td>
<td>1.4</td>
</tr>
<tr>
<td>FEV₁/Ht² (l/m²)</td>
<td>9.5</td>
<td>4.5</td>
</tr>
</tbody>
</table>
only to the FEV1/HT2 slope. It is highly probable that the acceleration of annual decline in FEV1 by smoking reflects obstructive changes in the respiratory system. In the case of pre-existing pulmonary fibrosis, however, the progression of restrictive disorders might be, at least in part, responsible for the acceleration, although its contribution to the FEV1/HT2 slope was about twofold larger than that of smoking.

In conclusion, no definite evidence was obtained that supports the possibility of an interaction of pre-existing mild pulmonary fibrosis and low exposure to chrysotile asbestos on the accelerated annual decline of FEV1, and FVC. Even if it is mild and stable, however, pre-existing pulmonary fibrosis is an independent risk factor for accelerated annual decline of FEV1, irrespective of exposure. As this study was based on only a few subjects, and past exposure before the start of the study was not considered, further investigation is needed to confirm the results.

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