Relation between exposure to respirable silica dust and silicosis in a tungsten mine in China

D Pang, S C Fu, G C Yang

Abstract
To estimate the quantitative relation between exposure to respirable silica dust and risk of an attack of silicosis, 1151 workers exposed to silica dust and employed from 1958 to 1987 in a tungsten mine in China were investigated. The results showed that the ratio of respirable silica dust concentration to total silica dust concentration was 0·529. Then, the total silica dust concentration in historical surveillance and monitoring data was converted to respirable silica dust concentration. The free silica content in respirable dust determined by x-ray diffraction averaged 24·7%. Multiple logistic regression was used for the dichotomous dependent variables (presence or absence of silicosis). The independent variables in the multiple logistic regression with presence of silicosis as the dependent variable were age when first exposed, tuberculosis (presence or absence), and cumulative exposure to respirable silica dust. The partial regression coefficient of individual cumulative exposure was estimated as 0·079. It implied a positive association between exposure to respirable silica dust and risk of an attack of silicosis. The exposure limit for respirable silica dust was estimated as 0·24 mg/m³ under given conditions.

Respirable dust is defined as that reaching the alveoli and is assumed to be associated more with pneumoconiosis than is total dust. As a result, health standards for respirable dust have been introduced by many countries—for example, America, Japan, Australia—in the context of prevention of pneumoconiosis. We are presently conducting a retrospective epidemiological study of selected workers from a tungsten mine in China to establish the quantitative relation between exposure to respirable silica dust and silicosis. The aim of the present study is to provide epidemiological evidence for setting and revision of the exposure limit for respirable silica dust.

Subjects and methods
Subjects
The study took advantage of both historical surveillance and monitoring data collected in the period from 1958 to 1987.

Men eligible for inclusion in the study were underground miners hired from 1958 to 1987 inclusive, without exposure to dust before 1958. Those who were previously exposed to other kinds of dusts, such as tungsten, were excluded. The criteria for inclusion were met by 1151 men.

Exposure and estimation
Information about the mine and the estimation of individual exposures by extrapolation based on fixed point membrane filter measurement has been published elsewhere. The total and respirable silica dust concentrations at workplaces of major job titles were measured. The free silica content in respirable dust was determined by x-ray diffraction.

Statistical methods
The relation between respirable dust concentration and total dust concentration was assessed by a linear regression model. The relation between silicosis and cumulative respirable silica dust concentration was determined by multiple logistic linear regression with the logit procedure of the SAS package.¹

Quality control measures
Quality control measures were instituted to ensure the quality of information collected. All data were checked when completed to establish that all items had been attempted. When completed all data were checked by computer procedures to ensure no logistic mistakes. Then all data were checked manually to ensure no entry errors. An airflow calibrating meter was used to calibrate the flow of samplers.

Results
Respirable and total dust concentrations
The figure shows the correlation between respirable and total dust concentrations at workplaces in the underground mine. A linear trend was found. Table
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Regression of respirable and total dust concentrations.

1 shows the t test statistics of regression. The regression coefficient was 0.529 (p < 0.001). It implied that the respirable dust concentration correlated with the total dust concentration. According to the regression coefficient, the ratio of respirable dust concentration to total dust concentration was 0.529. Therefore the respirable dust concentration was estimated based on total dust concentration in historical surveillance and monitoring data.

FREE SILICA CONTENT OF RESPIRABLE DUST
The arithmetic mean of six samples was 24.7%.

RESPIRABLE SILICA DUST CONCENTRATION AND SILICOSIS
Table 2 presents the results of the multiple logistic regression with presence of silicosis as the outcome. The partial regression coefficient between the cumulative exposure to respirable silica dust and silicosis was statistically significant (β = 0.079, p < 0.05) after allowing for the age of first exposure and tuberculosis (TB). The probability of developing silicosis correlated positively with cumulative exposure to respirable silica dust. The multiple logistic regression model can be expressed as follows:

\[ p = \frac{1}{1 + \exp(-((-7.331) + 0.079CUMCON + 0.108AGE1 + 1.59TB)))} \]

\[ CUMCON = C \times U \]

where:
- \( p \) = The probability level of developing silicosis.
- \( CUMCON \) = Cumulative exposure to respirable silica dust in units of mg years/m³.
- \( AGE1 \) = Age of first exposure (y).
- \( TB \) = Tuberculosis before silicosis developed.
- \( C \) = Respirable concentration (mg/m³).
- \( U \) = Duration of exposure (y).

If the sensitivity of the model was 95%, a probability level for classification of silicosis of 0.8% was derived. An exposure limit of 0.24 mg/m³ was estimated according to equations (1) and (2) when given a probability level for classification of silicosis as 0.8%, a duration of exposure of 30 years, and TB as 0 (those without TB).

Discussion
Silicosis is a chronic occupational disease with a long period of latency and it is important to establish its relation to respirable dust. To determine respirable dust concentrations it is necessary to convert total dust concentration in historical surveillance and monitoring data to respirable dust concentration by making experiments on respirable and total dust concentrations. Our results showed that the ratio of respirable to total dust concentrations was 0.529. It was assumed that changes in the ratio were small over various periods because working conditions and dust protective measures were stable. Estimates of respirable dust concentration were therefore derived from the total dust concentrations by applying a conversion factor of 0.529.

The probability of developing silicosis was positively correlated with cumulative exposure to respirable silica dust (table 2). This would be consistent with published evidence. The less the sensitivity of the regression model, the better the protection of workers from silicosis. It was accepted that the sensitivity was 95%. The age of first exposure was assumed to be 18 years according to the 12th clause of the Act of Silicosis of the People’s Republic of China, which stated that workers were “. . . not eligible to participate in dust work without being more than 18 years old.” The duration of exposure was assumed as 30 years taking the duration of employment in a worker’s whole life as 30–40 years. The free silica content in respirable silica dust was 24.7%. A respirable dust concentration of 0.24 mg/m³ was estimated based on these conditions. We conclude that the exposure limit of 0.24 mg/m³ would be safe allowing for epidemiological evidence.

### Table 1 Regression analysis of respirable and total dust concentrations

<table>
<thead>
<tr>
<th>Independent variable</th>
<th>Dependent variable</th>
<th>No of samples</th>
<th>t Value</th>
<th>p Value (one sided)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Respirable dust</td>
<td>Total dust</td>
<td>25</td>
<td>15.247</td>
<td>0.000</td>
</tr>
</tbody>
</table>

### Table 2 Linear multiple logistic regression with presence of silicosis as outcome and with age of first exposure, TB, and cumulative exposure to respirable silica dust as independent variables

<table>
<thead>
<tr>
<th>Variable</th>
<th>β (SD)</th>
<th>( \chi^2 )</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>-7.331 (0.877)</td>
<td>69.95</td>
<td>0.0001</td>
</tr>
<tr>
<td>Cumulative concentration</td>
<td>0.079 (0.031)</td>
<td>6.38</td>
<td>0.0115</td>
</tr>
<tr>
<td>Age of first exposure</td>
<td>0.108 (0.029)</td>
<td>13.71</td>
<td>0.0002</td>
</tr>
<tr>
<td>Tuberculosis</td>
<td>1.59 (0.609)</td>
<td>0.83</td>
<td>0.0089</td>
</tr>
</tbody>
</table>

Regression of respirable and total dust concentrations.

![Graph showing the relationship between respirable dust concentration and total dust concentration.](image)

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Inhalation Fever: A proposed unifying term for febrile reactions to inhalation of noxious substances

Over the past several years there have been many publications that consider respiratory illnesses of farmers. Much of the interest has centred around the description of a febrile illness caused by exposure to organic dust that has been called several names including: grain fever, pulmonary mycotoxicosis, silo unloader’s syndrome, and organic dust toxic syndrome (ODTS). There is now sufficient evidence to suggest that these febrile attacks are separate from allergic alveolitis. The physiological impact of the febrile illness on the lung is minimal whereas in allergic alveolitis one sees abnormal blood gases, chest radiographs, and pulmonary function tests.

In the modern industrial era there has been extensive writing about febrile responses to the inhalation of metal fumes and dusts such as metal fume fever among welders and foundry workers. As plastics emerged papers appeared describing febrile illnesses in those who work with complex polymers.

The clinical picture of these inhalation ailments is similar. All have a latent period of several hours between the exposure and the onset of fevers, chills, malaise, myalgias, and chest tightness. Patients often have a leucocytosis with a left shift. There are reports of chemical pneumonitis and pulmonary oedema in some metal and plastic exposures and a single report suggesting more severe lung involvement after exposure to organic dust in an agricultural setting, but most reports suggest that little lung damage occurs.

We are struck by the described similarities and believe that a generic term is best for this family of respiratory ailments. In the absence of a single aetiological agent for these very different exposures we think that the fevers and symptoms represent a common pathway in which the lung reacts to noxious substances. We suggest that they all be called inhalation fever, a term originating from 1978. This avoids the colourful but confusing litany of names currently in use. Henceforth we propose that organic dust toxic syndrome, all the fume fevers, spelter shakes, mill fever, Monday fever, card room fever, and brass founders ague etc, be called simply Inhalation fever.

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