Airway responsiveness, respiratory symptoms, and exposures to soluble oil mist in mechanical workers

N Massin, A B Bohadana, P Wild, P Goutet, H Kirstetter, J P Toamain

Abstract

**Objectives**—To assess the relation between measured levels of exposure to soluble oil mists in a plant manufacturing ball bearings, and both respiratory symptoms and airway responsiveness in the workforce.

**Methods**—114 male workers exposed to oil mist and 55 unexposed male controls from nearby factories were studied. Soluble oil mist concentrations were measured with area samplers. Respiratory symptoms were assessed by questionnaire and measurement of airway responsiveness to methacholine with an abbreviated method. Subjects were labelled positive to methacholine airway challenge (MAC+) if their forced expiratory volume in one second (FEV1) fell by ≥20%. The linear dose-response slope was calculated as the percentage fall in FEV1 at the last dose divided by the total dose given.

**Results**—Geometric mean concentrations of oil mists ranged from 0.65 mg/m3 (GSD = 1.29) to 2.20 mg/m3 (GSD = 1.55) based on 92 measurements obtained from 1979–93. The prevalence of chronic cough or phlegm, bouts of bronchitis, and dyspnoea was greater among exposed workers than among controls (odds ratio (OR) 4.64, P = 0.002 for chronic cough and phlegm). After adjustment for smoking and age, dyspnoea was significantly related to an index of cumulative exposure to oil mist (OR 1.44, P = 0.006/10 y.mg/m3). The proportion of MAC+ subjects was similar in the two groups. However, after adjustment for baseline FEV1 and age, the dose-response slope was significantly steeper among exposed workers than among controls (P = 0.01), a finding indicating airway hyperresponsiveness in the exposed workers. Furthermore, the dose-response slope was significantly related to baseline FEV1, age, and, after adjustment for FEV1, the index of cumulative exposure to oil (P = 0.004).

**Conclusion**—Subjects with exposure to soluble oil mist in the metal industry are at risk of developing both respiratory symptoms and airway hyperresponsiveness.

(Occ Environ Med 1996;53:748–752)

Keywords: airway responsiveness; respiratory symptoms; soluble mineral oils

Worldwide, several million workers are regularly exposed to mists of machining fluids in the metal industry. These fluids are complex mixtures of oil (mineral or synthetic) and other substances—for example, water, emulsifiers, corrosion inhibitors, bactericides, etc—which are used mainly as lubricants and coolers to prevent the welding of the metal parts to the cutting tool itself. Previous investigations have shown adverse respiratory effects of occupational exposure to oil mists: epidemiological studies showed that workers exposed to various types of mineral oil mists were more likely to have increased rates of symptoms of chronic bronchitis than control subjects.

Also, several patients exposed to oil mists had occupational asthma and were referred to a respiratory clinic because of asthmatic symptoms. Finally, a recent report showed a significant decline in forced expiratory volume in one second (FEV1) during shifts among non-asthmatic machine operators exposed to relatively low concentrations of mists of machining fluids in the metal industry.

Over the past two decades, the proportion of oil in the composition of machining fluids fell drastically from 100% in straight cutting oils to ≤10% nowadays in soluble mineral oil. In this study we consider the question of the relation between levels of exposure to soluble oil mists and respiratory symptoms and indices of airway responsiveness to methacholine, a matter on which there is only limited information. The results were compared with those obtained in a group of unexposed control subjects.

**Materials and methods**

**STUDY DESIGN AND SUBJECTS**

A cross-sectional survey of 114 male employees was carried out, representing 85% of the workforce at a plant manufacturing ball bearings in eastern France which has used only soluble mineral oil over the past 25 years. All workers have been chronically exposed to mists of soluble mineral oils for at least one year. The control population was a group of 55 blue collar male workers who had never been at occupational risk of exposure to mineral oils or any other respiratory hazard. They represented 90.2% of the workforce (n = 61) at two manufacturing plants: a food distribution plant (n = 27) and a stationery factory (n = 28). They were from the same geographical region and worked the same shifts as the exposed workers; both groups were examined in the autumn of 1993. Written and informed consent was obtained from all workers.
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consent was given by all subjects. Information was obtained about age, working hours a day, long term duration of work, and occupational history. Table 1 shows the prestudy characteristics of exposed workers and controls.

EXPOSURE MONITORING

The current concentrations of soluble mineral oils in the workplaces were measured over periods of two to three hours with area samplers. Ambient air was aspirated through glass fibre filter paper (GF/C Whatman, 37 mm in diameter) with small vacuum pumps (Dupont S-2500, Kennett Square, PA, USA) the flow rate of which was kept at 1-0 l/min. The oil was extracted from the filter with dichloromethane and its concentration was determined by gravimetric analysis. Repeated samples were obtained from 1979 to 1993 to take changes in the working conditions into account. It has to be stressed that this methodology was developed to assess straight cutting oils consisting mainly of mineral oil. In fact, the values measured in this study can only be considered as an indicator of the exposure as <10% of the mist is soluble in dichloromethane.

For a given area, the concentration of oil mist was calculated as the geometric mean value of the measurements obtained in that area: the figures were rounded to the nearest whole number to provide a score which was assigned individually to all workers in the area. Next, for a given worker, the exposure score was multiplied by the number of working months in the area for the period of employment as a whole. This allowed for the calculation of an index of estimated exposure to oil mist—namely, the cumulative exposure to oil.

RESPIRATORY HEALTH STATUS

Medical history

Detailed histories of respiratory diseases and smoking habits were recorded. All subjects completed a modified version of the European Coal and Steel Community questionnaire on respiratory symptoms which was given by the same experienced interviewer. The questionnaire emphasised the past and present personal and family histories of cough, phlegm, or both, asthma, wheeze, and dyspnoea. Chronic bronchitis was defined as cough and phlegm for at least three months each year for not less than two successive years. The term chronic cough or phlegm was used to define subjects complaining of cough, or phlegm, or both regardless of duration: thus it encompassed those subjects who fulfilled the criteria for chronic bronchitis and those who did not. Bouts of bronchitis was the term coined to define those subjects who answered yes to the question: “Have you ever experienced an increase in the severity of cough and in the volume of sputum production of at least three weeks in duration during the past three years?” Dyspnoea on exertion was considered to be present when the subjects complained of breathlessness when walking up a slight hill. Asthma was considered to be present if the subjects answered yes to the question: “Have you ever been diagnosed as having bronchial asthma?”

Non-smokers were defined as subjects who had never regularly smoked one or more cigarettes a day or had smoked one or more cigarettes a day for less than one year. Current smokers were defined as subjects who reported regular smoking of one or more cigarettes a day for at least one year. Ex-smokers were subjects who reported smoking one or more cigarettes regularly in the past but who had stopped smoking at least one year before the study.

Pulmonary function tests

Spirometry was carried out by the same experienced technician (JPT) with an electronic thermister spirometer (Spiro-Analyzer ST 300, Fukuda Sangyo, Tokyo, Japan). The following indices were obtained by having the subject expire forcefully and maximally after a maximal inspiratory manoeuvre: forced vital capacity (FVC), forced expiratory volume in one second (FEV1), forced expiratory flow between 25 and 75% of the FVC (FEF25-75), and maximal expiratory flows at 25%, and 50% of the FVC (Vmax25, Vmax50, respectively). At baseline each subject performed at least three reproducible forced expiratory manoeuvres (within 5% for FVC and FEV1); thereafter only two reproducible curves were required. The curve with the highest sum of FVC + FEV1 was used for statistical analysis. The results were expressed as standardised residuals (SR) (SR = (observed value—predicted value)/residual standard deviation (RSD)) as recommended by a committee of the European Community for Steel and Coal. An SR of zero indicates that the observed value is equal to the control value (at the 50th percentile); SRs of −1.64 or 1.96 indicate that the results are at the 5th percentile and the 97.5th percentile respectively.

Airway responsiveness

As the examinations were carried out during workshifts, an abbreviated version of the methacholine airway challenge (MAC) test was used. Three cumulative doses of methacholine (0.5 μmol, 2.5 μmol, and 7.5 μmol—that is, 100 μg, 500 μg, and 1500 μg respectively) were given with a nebuliser (Mediprom FDC 88—Paris, France) delivering doses of 0.5 μmol methacholine per breath. A noseclip was worn and the aerosol

Table 1 Anthropometric characteristics, smoking history, and duration of exposure in workers exposed to soluble mineral oil mist and controls

<table>
<thead>
<tr>
<th></th>
<th>Exposed workers (n = 114)</th>
<th>Controls (n = 55)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (mean (SD), y)</td>
<td>38 (9)</td>
<td>30 (8)</td>
</tr>
<tr>
<td>Weight (mean (SD), kg)</td>
<td>81 (12)</td>
<td>73 (10)</td>
</tr>
<tr>
<td>Height (mean (SD), cm)</td>
<td>176 (8)</td>
<td>174 (6)</td>
</tr>
<tr>
<td>Smoking status:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Current smokers (n (%))</td>
<td>47 (41)</td>
<td>37 (37)</td>
</tr>
<tr>
<td>Ex-smokers (n (%))</td>
<td>41 (36)</td>
<td>0</td>
</tr>
<tr>
<td>Never smokers (n (%))</td>
<td>26 (23)</td>
<td>18 (33)</td>
</tr>
<tr>
<td>Smokers (mean (SD), pack-years)</td>
<td>12 (9)</td>
<td></td>
</tr>
<tr>
<td>Ex-smokers (mean (SD), pack-years)</td>
<td>12 (11)</td>
<td></td>
</tr>
<tr>
<td>Duration of exposure (mean (SD), y)</td>
<td>15 (8)</td>
<td></td>
</tr>
</tbody>
</table>
inhaled through the mouth slowly. Then the breath was held for five seconds. Spirometry was performed in the sitting position, before and three minutes after the inhalations of methacholine. The challenge test was discontinued either after the inhalation of the third dose of methacholine or if the FEV₁ fell by \( \geq 20\% \) below the baseline value.

Subjects who experienced a fall in FEV₁ of \( \geq 20\% \) were classified as having a positive MAC test (MAC⁺)—that is, the airway was hyperresponsive. A linear dose-response slope was calculated by the method proposed by O'Connor and colleagues \(^\text{13}\) as the percentage fall in FEV₁, at the last dose divided by the total dose of methacholine given.

**ETHICS**

The study was approved by the local medical ethics committee.

**STATISTICAL ANALYSIS**

To be able to apply a multivariate analysis to the dose-response slope, a transformation to normalise the data was applied. The transformation \( 1/(\text{slope} + 2.5) \) in the family of shifted logarithmic and shifted inverse transformations was found to be optimal for a large unexposed population. \(^\text{8}\)

Statistical analysis was carried out with the SAS statistical software. \(^\text{14}\) Multiple logistic regression analyses were used to assess the effect of the exposure on the symptoms while adjusting for smoking status and age (for dyspnoea).

Multiple linear regression was used to describe the effect of exposure on the baseline spirometric variables (expressed as SRs) adjusted on tobacco consumption, and on the transformed dose-response slope adjusted for baseline FEV₁, and age. We did not include smoking in this model as we found that this variable was unrelated to bronchial responsiveness. The stability of the variance and approximate linearity in the linear models were checked on residual plots.

**RESULTS**

**EXPOSURE ASSESSMENT**

Overall, 92 samplings were performed: 20 in the cutting area and 72 in the machining area. In the cutting area, the geometric mean concentration of measurements carried out in 1989 was 1.49 (GSD 1.22) mg/m³; as no major hygiene measures aimed at improving the air quality were carried out in this area between 1979 and 1993, this value was considered to be representative of the working conditions in this period. In the machining area, the geometric mean concentration value of 46 samplings performed between 1979 and 1989 was 2.20 (GSD 1.55) mg/m³. After hygiene measures aimed at improving ventilation were adopted in 1990, the geometric mean value of 26 measurements decreased to 0.65 (GSD 1.29) mg/m³. Both these mean values were taken into account in the exposure assessment of subjects working in this area.

**RESPIRATORY SYMPTOMS AND BASELINE LEVEL OF PULMONARY FUNCTION**

Overall, the prevalence of respiratory symptoms tended to be greater among the exposed workers than among unexposed controls. The proportion of subjects with chronic bronchitis, chronic cough or phlegm, or bouts of bronchitis was three to nine times as great among workers exposed to oil mist than in unexposed controls; for chronic cough or phlegm the differences were significant (table 2). After adjustment for smoking and age (on average exposed workers were eight years older than controls), dyspnoea seemed to be significantly related to the index of cumulative exposure to oil mist, but for asthma the differences were not significant (table 2). It is interesting to notice, however, that five out of six exposed workers developed asthma after they were hired, whereas both controls with asthma had had it since childhood: in one asthma persisted through adult life and in the other it had disappeared by the age of 11.

After adjustment for tobacco consumption, workers exposed to oil mist had baseline pulmonary function variables similar to those of unexposed controls (table 3). After adjustment for tobacco consumption, baseline FEV₁ was not related to oil exposure.

**AIRWAY RESPONSIVENESS**

The methacholine airway challenge test was carried out in all exposed workers (n = 114) and unexposed controls (n = 55). A positive MAC test (MAC⁺ = FEV₁ fall \( \geq 20\% \)) was found in 14 out of the 169 subjects tested; the proportion in exposed workers was similar to that in controls (table 4). In contrast, mean dose-response slope values were significantly lower (steeper slope) in smokers than non-smokers.
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Table 4 Airway responsiveness to methacholine among workers exposed to soluble mineral oil mists and controls

<table>
<thead>
<tr>
<th>Factor</th>
<th>Exposed workers (n = 114)</th>
<th>Controls (n = 55)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>MAC + (%), n (%)</td>
<td>10 (9)</td>
<td>4 (8)</td>
<td>&gt; 0.5*</td>
</tr>
<tr>
<td>Dose-response slope (1/(slope + 2.5)) (mean (SD))</td>
<td>0.27 (0-08)</td>
<td>0.30 (0-07)</td>
<td>0.03†</td>
</tr>
</tbody>
</table>

*P* test.
†Based on a linear model after adjusting for baseline FEV₁ and age.
MAC = positive spirometric methacholine airway challenge (% fall in FEV₁, by > 20%); dose-response slope = (slope = % fall in FEV₁/mmol methacholine).

Table 5 Multiple regression of the dose-response slope* on baseline FEV₁, age, and exposure to soluble oil mists expressed in terms of cumulative exposure

<table>
<thead>
<tr>
<th>Factor</th>
<th>Coefficient</th>
<th>SE</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Constant</td>
<td>0.05256</td>
<td>0.0086</td>
<td>0.0001</td>
</tr>
<tr>
<td>Baseline FEV₁(l)</td>
<td>0.004269</td>
<td>0.0008</td>
<td>0.029</td>
</tr>
<tr>
<td>Age (y)</td>
<td>0.00195</td>
<td>0.0003</td>
<td>0.004</td>
</tr>
<tr>
<td>Cumulative exposure (10 y.mg/m²)</td>
<td>-0.00999</td>
<td>0.0003</td>
<td>0.004</td>
</tr>
</tbody>
</table>

*Dose-response slope = 1/(% fall in FEV₁/mmol + 2.5).

workers exposed to oil than in controls, a finding indicating overall airway hyperresponsiveness in the exposed group (table 4).

Multiple regression analysis was used to assess the relative contribution of measured variables to dose-response slope values. The only variables likely to influence dose-response slope values were baseline FEV₁, age, and exposure to soluble oil mist. Table 5 shows the significance of these variables included in the model for the group as a whole. From this model it can be seen that 10 years of exposure at 1 mg/m² is modelled to have the same effect on dose-response slope as a 230 ml decrease in the baseline FEV₁.

Discussion

This study shows that workers exposed to soluble mineral oil mist have more respiratory symptoms and are significantly more hyperresponsive to methacholine than are comparable unexposed controls.

To the best of our knowledge, the recent paper by Ameline and colleagues is the only investigation published in English dealing with airway responsiveness to methacholine in workers exposed to soluble oil mist. These authors examined the prevalence of respiratory symptoms, ventilatory abnormalities, and airway hyperresponsiveness in workers exposed to mists of straight cutting oil (n = 40), soluble mineral oil (n = 51), or a mixture of straight cutting oil and soluble mineral oil (n = 139), and compared the results with those found in a group of unexposed controls (n = 78). Overall, only slight adverse effects on respiratory symptoms and lung function were found which were restricted to workers exposed to straight cutting oil. Moreover, no effect of exposure was found on airway responsiveness, whatever group was considered.

Theoretically, the differences in the degree of airway responsiveness of our workers compared with those exposed to soluble mineral oil in the study of Ameline and colleagues could be explained by various factors including the exposure, the challenge test, or both. It is unlikely that exposure was the only factor involved as the type of oil used as well as the mean duration of exposure were similar in the two studies. Differences related to the technique of giving the methacholine can also be discarded confidently as, in both studies, similar doses of this agent were given through identical nebulisers, and the test results were expressed with similar indices.

A more likely source of disagreement lies in the way of calculating the dose-response slope: indeed, whereas we added a constant of +2.5 (see methods section) to our slopes to achieve a distribution close to normal as possible, Ameline and colleagues chose to omit negative slopes. However, when they reanalysed their data (for workers exposed to soluble oil mist) with our methods, their results became similar to ours (P Wild, personal communication).

The significance of non-specific airway hyperresponsiveness in occupational samples in general is not completely understood. In other types of occupational exposure airway hyperresponsiveness is often associated with respiratory symptoms and impaired lung function. Although such cross-sectional associations do not indicate whether airway hyperresponsiveness itself is a risk factor that precedes the development of these abnormalities or not, their documentation is important for they provide the rationale for longitudinal studies necessary to answer this question. In this study we show that workers exposed to oil mist present airway hyperresponsiveness to methacholine which is significantly related to an index of cumulative exposure to oil. We also showed that these workers have significantly increased rates of chronic cough or phlegm and dyspnoea. Dyspnoea was related to an index of cumulative exposure to oil mist.

In our opinion, although these associations do not constitute formal evidence of a causal relation they surely provide a strong argument pointing in that direction. Considering the current concept on the mechanisms of non-specific airway hyperresponsiveness we are tempted to suggest that in our workers this resulted from airway inflammation due to longstanding exposure to oil mist. Theoretically, such sensitisation might have resulted either from airway sensitisation to specific components of oil mists or to a non-specific irritant airway effect, or both. Indeed, the sensitisation to specific components of oil mists has been documented in cases of occupational asthma whereas the non-specific irritant airway effect has been used to explain acute lung response to sustained exposure to oil mist.

This study is limited by the fact that the exact composition of the soluble mineral oil mists our workers were exposed to is unknown, thus preventing further consideration of the possible agent or agents likely to be involved in a theoretical inflammatory process. However, this by no means reduces the importance of the impairment in respiratory health that we found in workers exposed to soluble oil mist.

To summarise, this study shows that workers exposed to soluble oil mist have significantly
more airway hyperresponsiveness to methacholine and chronic respiratory symptoms than unexposed controls. Careful controlled longitudinal studies are necessary to find whether or not this abnormality represents an early manifestation of occupational asthma or chronic airway obstruction.

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*Occup Environ Med* 1996 53: 748-752
doi: 10.1136/oem.53.11.748

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