The effect of insecticide aerosols on lung function, airway responsiveness and symptoms in asthmatic subjects

C.M. Salome*, G.B. Marks*#, P. Savides*, W. Xuan*, A.J. Woolcock*


ABSTRACT: The object of this study was to compare the effect of standard and "low irritant" insecticide aerosols on lung function, airway hyperresponsiveness (AHR) and symptoms in asthmatic subjects.

A double blind randomized, crossover study was conducted in 25 asthmatic subjects who reported sensitivity to insecticide aerosols. All subjects were exposed for 30 min, on separate occasions, to two standard insecticide formulations (A and B), one low irritant formulation (C) and a negative control aerosol. Spirometric function and chest, nose and eye symptoms were recorded during, and for 90 min after, the exposure. AHR to methacholine was measured 90 min after the exposure. Compared to the negative control, the maximum fall in forced expiratory volume in one second (FEV1) was slightly greater after standard insecticides (mean difference from control=95% confidence interval: aerosol A, 3.3±3.6%, p=0.08; aerosol B, 5.1±4.7%, p=0.04). AHR was significantly more severe (mean difference from control: aerosol A, 0.35±0.29 doubling doses, p=0.028; aerosol B, 0.52±0.43 doubling doses, p<0.028), and symptoms were more severe. The low irritant test aerosol (C) did not differ significantly from the negative control with respect to FEV1, AHR or symptoms.

It is concluded that some insecticide aerosols trigger symptoms and falls in lung function in some people with asthma. Furthermore, these aerosols may also increase airway hyperresponsiveness, although the mechanism of this effect has not been determined. The low irritant formulation did not appear to have the same effects.


Patients with asthma commonly report exposure to aerosol insecticides as a triggering factor for asthma symptoms. Pyrethrum and synthetic pyrethroids have been used as insecticides for most of this century and their use in aerosol form is now widespread in the developed world. From as early as 1930 there have been reports of asthma attacks triggered by pyrethrum [1], and more recently there has been a report of a death due to sudden bronchospasm following inhalation of a pyrethrin shampoo [2]. However, there has been only one previous systematic study of the role of pyrethrum insecticides in asthma [3]. This study showed that insecticide aerosols can cause symptoms and changes in lung function in some asthmatic subjects. However, the sample size in this study was small, and only one subject had a clinically relevant fall in forced expiratory volume in one second (FEV1).

Because of a perceived need for special formulations for asthmatic and sensitive subjects, insecticide manufacturers have focused on developing new low irritant insecticide formulations. Claims that such formulations are particularly safe for asthmatics are widely promoted, but have received no formal objective testing in asthmatic subjects.

The present study was undertaken to determine whether insecticide aerosols have any effect on lung function, airway hyperresponsiveness or symptoms in subjects with a history of symptoms on exposure to domestic fly-sprays. Furthermore, the authors wished to determine whether a "low irritant" insecticide formulation had any adverse effects.

Methods

Subjects

Twenty-five asthmatic subjects, aged >16 yrs, with a previous history of sensitivity to insecticide aerosols were recruited from the Asthma Clinic at Liverpool Hospital (Sydney, Australia). All subjects had a previous diagnosis of asthma, required regular therapy, and had existing airway hyperresponsiveness to methacholine, with a provocative dose causing a 20% fall in FEV1 (PD20) <4 µmol and baseline FEV1 >60% predicted normal. Subjects were excluded from the study if they had other major illnesses, were pregnant or breast feeding, or had had a serious exacerbation of their asthma, requiring an urgent doctor or hospital visit in the last 4 weeks.

The protocol for the study was approved by the Ethics Review Committee of the South Western Sydney Area Health Service (Australia).

Study design

The study was a randomized, double-blind cross-over study of the effects of insecticide aerosols on people with asthma who reported a history of symptoms on exposure to...
fly-sprays. Subjects were exposed to the insecticide aerosols on four separate days, at least 1 week apart. All subjects were exposed to three of the aerosols. These were a standard formulation, which had been shown to cause symptoms in a previous study (aerosol A) [3], a low irritant formulation (aerosol C) and a negative control aerosol. Because the older standard formulation is no longer commercially available, a fourth exposure was used to determine the effects of two other standard insecticide formulations (aerosols B1 and B2), both of which are currently commercially available. Subjects were randomly divided into two groups and exposed to one or other of the two sprays. The contents of all test aerosols are shown in table 1. The aerosols were activated to give a starting dose which approximates a real life situation, followed by a second dose comparable to the doses used in the study by Newton and Breslin [3].

The effects of the exposure were determined by examining changes in symptoms affecting the chest, nose and eyes, as well as changes in spirometric function and airway responsiveness to methacholine. Spirometric function and symptoms were assessed at 5 and 15 min after each dose, and then at 15, 30, 60 and 90 min after leaving the exposure chamber. Airway responsiveness was measured 90 min after the end of each exposure.

### Aerosol exposure

An exposure chamber was developed from a room at the Asthma Clinic at Liverpool Hospital. The room contained a desk and chair and was equipped with appropriate exhaust systems to adequately ventilate the room. Surfaces in the room were covered during the aerosol exposures. For each of the test aerosols two dose levels were assessed. The test aerosols were activated within the exposure chamber, firstly, for 4 s, and then, 15 min later, for 32 s. Based on the volume of the chamber (23 m$^3$) and the output of the aerosol devices (1.7 g s$^{-1}$, of which 0.35% is active natural pyrethrum), it is estimated that the maximum concentration of the active pyrethrum was 8.3 mg m$^{-3}$ at the time of the 32 s dose. The maximum exposure is comparable to that which caused symptoms in the study by Newton and Breslin [3], but is less than half the maximum dose used in their study. The estimated exposure concentration in this study was less than one tenth the "No observable effect level" (100 mg m$^{-3}$) and was close to the threshold limit value (5 mg m$^{-3}$) [4–6].

### Table 1. Contents of insecticide aerosols tested

<table>
<thead>
<tr>
<th>Aerosol sample</th>
<th>Active insecticide</th>
<th>% w/w</th>
<th>Propellant Base</th>
</tr>
</thead>
<tbody>
<tr>
<td>Negative control</td>
<td>-</td>
<td></td>
<td>HC Water</td>
</tr>
<tr>
<td>A Standard insecticide-older formulation</td>
<td>Pyrethrins Tetramethrin Piperonyl butoxide N-Octyl bicycloheptene dicarboxamide</td>
<td>0.30 0.09 1.50</td>
<td>HC Solvent</td>
</tr>
<tr>
<td>B1 Standard insecticide-commercial aerosol</td>
<td>Allethrin Tetramethrin Piperonyl butoxide N-Octyl bicycloheptene dicarboxamide Pyrethrins</td>
<td>0.75 0.27 0.24 0.41</td>
<td>HC Solvent</td>
</tr>
<tr>
<td>B2 Standard insecticide-commercial aerosol</td>
<td>Pyrethrins Piperonyl Butoxide</td>
<td>0.66 0.35 1.61</td>
<td>HC Water</td>
</tr>
<tr>
<td>C Low irritant aerosol</td>
<td>Bioallethrin Bioresmethrin</td>
<td>0.28 0.05</td>
<td>HC Water</td>
</tr>
</tbody>
</table>

%w/w: the weight of the constituent chemical as a percentage of the weight of the insecticide formulation; HC: hydrocarbon.

### Spirometric function

Spirometric function was recorded on an electronic spirometer, with the subject standing. FEV1 and forced vital capacity (FVC) were measured and the highest of two measurements, reproducible to within 5% or 100 mL, were recorded. Per cent predicted values [7] were calculated.

### Airway responsiveness to methacholine

Airway responsiveness to methacholine was measured using the method of Yan et al. [8]. Methacholine was administered to the subjects via a hand-held De Vilbiss No.45 nebulizer (De Vilbiss Co., Somerset, PA, USA) in doubling doses, ranging 0.05–12.8 μmol, until the FEV1 fell by 20%. Response to challenge was measured by the dose response ratio, which is calculated as per cent fall FEV1/dose (μmol) [9, 10]. Short acting β-agonists were withheld for 6 h and long-acting β-agonists for 24 h prior to testing.

### Symptom scoring

Symptoms were assessed to determine both the type and the severity of symptoms. Chest symptoms were recorded as difficulty in breathing, chest tightness and cough; nose symptoms were recorded as running, blocking and sneezing; eye symptoms were recorded as itchy eyes, sore eyes and tears. A visual analogue scale (VAS) was used to measure the severity of the symptoms. For each of these nine symptoms, subjects were shown a 10 cm line, marked at one end "none" and at the other "the worst I can imagine" and asked to mark the point on the line which corresponded to the severity of the sensation they were experiencing at that moment. The length of the line, from zero ("none") to the point marked was measured in millimetres and used as an indicator of severity.

### Data analysis

**Lung function.** Changes in FEV1 and FVC during and after the exposure were calculated as per cent change from the baseline measured immediately prior to exposure. The proportion of subjects with a fall in FEV1 >15% during the exposure were calculated, and a comparison made between the test aerosol and the positive and negative controls, using...
McNemars test for proportions. The maximum per cent change in FEV1 for each subject during each exposure was calculated and compared by analysis of variance for repeated measures. The distribution of the maximum percentage falls in FEV1 did not differ significantly from the normal distribution, using the Kolmogorov-Smirnov statistic, following any of the exposures. Summary data for FEV1 are shown as means and 95% confidence intervals throughout this article.

Airway responsiveness. Values for dose response ratio (DRR) were log-transformed prior to statistical analysis and geometric mean DRR values on the four exposure days were calculated. The DRR values were compared by analysis of variance for repeated measures. The DRR values were adjusted for nonspecific responses to the negative control and a comparison made between the test aerosol, the positive control and the commercial insecticide aerosols. DRR values are reported as geometric means, and 95% confidence intervals. Changes in DRR are reported in terms of doubling dose difference with their 95% confidence intervals.

Symptoms. Scores for chest, nose and eye symptoms were calculated for each subject at each time point as the sum of the VAS scores recorded at each time point for the three chest symptoms, the three nose symptoms and the three eye symptoms. The maximum symptom score for each subject following each exposure was determined and the maximum change in symptom score was calculated by subtracting the score at baseline from the maximum score. The maximum changes in symptom scores after each exposure were compared by analysis of variance. In addition, to examine the time course of the changes in symptoms, mean symptom scores were calculated at each time point following each exposure. Symptom scores are reported as mean VAS scores and 95% confidence intervals.

Results
Symptom data were obtained from 25 asthma subjects with mean age 38±6 yrs. Lung function and bronchial challenge data from one subject were excluded because the subject was unable to perform reproducible expiratory manoeuvres. In the remaining 24 subjects baseline FEV1 was 80.8±5.4 % pred and baseline FVC was 80.0±5.4 % pred. All subjects had taken inhaled medication on most days in the month prior to the study, and 20 were taking regular inhaled corticosteroids. Two subjects were taking oral corticosteroids. On the basis of their airway hyperresponsiveness to methacholine, nine were classified as having severe or moderate airway hyperresponsiveness (AHR) and the remainder had mild AHR.

Thirteen subjects were exposed to aerosol B1 and 12 subjects to aerosol B2. The mean maximum fall in FEV1 was 9.7±5.0% after B1 and 9.2±7.0% after B2, and the mean increase in AHR from the negative control was 0.56 (-0.02–1.1) doubling doses after B1 and 0.47 (-0.26–1.2) doubling doses after B2. There were no significant differences between the effects of B1 and B2 aerosols on lung function, airway responsiveness and symptoms. Consequently, the data were combined and all subsequent analyses were performed on the combined data, and are shown below as aerosol B.

Lung function
Exposure to the insecticide aerosols had significant effects on lung function, and these effects differed significantly between the aerosols. Figure 1 shows the time course of mean falls in FEV1, with their 95% confidence intervals, for each of the four exposures. The fall in FEV1 was greatest at 20 min, five min after the 32 s activation, but it had returned to pre-exposure levels by the end of the observation period.

The maximum fall in FEV1 differed significantly between exposures (p=0.019) (table 2). Compared to the negative control the maximum fall in FEV1 was slightly greater after standard aerosol A, with a mean difference from negative control of 3.3±3.6% fall (p=0.08) and significantly greater after aerosol B, with mean difference of 5.1±4.7% fall (p=0.04). The maximum fall caused by aerosol C, the low irritant formulation, did not differ from the negative control, with a mean difference 1.3±3.2% fall (p=0.44). The fall in FEV1 caused by aerosol C was significantly lower than the response to aerosol B (p=0.039), and slightly lower than the response to aerosol A (p=0.078). The magnitude of the changes in FEV1 after the standard aerosols was not related to the baseline severity of AHR, measured after the negative control aerosol (for aerosol A, r=0.026 and for aerosol B, r=0.01).

The proportion of subjects with clinically relevant falls in FEV1 (falls ≥15%) after each of the aerosols is shown in table 2. The proportion was significantly greater after aerosol B than after the negative control (p=0.016), and was slightly greater after aerosol A than after the negative control (p=0.06). After the low irritant formulation the proportion of subjects with clinically relevant falls in FEV1 did not differ from the negative control (p=0.50).

Airway responsiveness
Airway responsiveness to methacholine was increased by exposure to the standard aerosols (table 2). Compared to the negative control, mean dose response ratio was significantly higher (more severe) after aerosol A (p=0.028) and aerosol B (p=0.028), but not after the low
irritant aerosol (p=0.60). These differences remained significant when the responses to the active aerosols were adjusted for the response to the negative control (p=0.0083) (fig. 2). The mean increase in responsiveness compared to the negative control was 0.35±0.29 doubling doses for aerosol A and 0.52±0.43 doubling doses for aerosol B.

Symptoms

Exposure to the insecticide aerosols caused increases in chest, nose and eye symptoms. The maximum effect occurred 5 min after the 32 s activation of each aerosol, but had returned to baseline levels by the end of the observation period. Figure 3 shows the time course of changes in the VAS scores for chest symptoms, nose symptoms and eye symptoms. The mean change in maximum VAS scores for chest symptoms, nose symptoms and eye symptoms are shown in table 2. Symptom scores tended to be highest after aerosol B and aerosol A, intermediate after aerosol C and lowest after the negative control. Compared to the negative control, aerosol A caused significantly more severe nose and eye symptoms, aerosol B caused significantly more severe chest and nose symptoms. Following aerosol B the magnitude of the maximum fall in FEV1 was significantly correlated with the maximum score for chest symptoms (r=0.43, p=0.02), but not with nose or eye symptoms. The low irritant aerosol did not differ significantly from the negative control for any set of symptoms as assessed by the multi-range hypothesis test.

Discussion

The authors have reported the results of a randomized, double-blind crossover study of the effects of insecticide aerosols on people with asthma who report a history of symptoms on exposure to fly-sprays. The study compared the effects of three standard insecticide formulations, one of which had previously been shown to cause symptoms [3], with those of a negative control. Compared to the negative control, aerosol A caused significantly more severe nose and eye symptoms, aerosol B caused significantly more severe chest and nose symptoms. Following aerosol B the magnitude of the maximum fall in FEV1 was significantly correlated with the maximum score for chest symptoms (r=0.43, p=0.02), but not with nose or eye symptoms. The low irritant aerosol did not differ significantly from the negative control for any set of symptoms as assessed by the multi-range hypothesis test.
asthmatic patients is not known, but clinical experience suggests that it may be common. In a community population sample, 22.7% of respondents reported chemical intolerance, defined as feeling ill on exposure to at least three of a panel of five common chemicals (paint, pesticides, car exhaust, new carpet and perfume) [11]. Among these subjects, asthma and respiratory symptoms, including wheeze, and chest tightness were common. For this study, subjects with a history of symptoms associated with insecticide use were selected because they were likely to be the subjects most at risk from an adverse effect of insecticide aerosols, and most likely to be targeted by marketing strategies aimed at insecticide sensitive asthmatics.

This analysis has confirmed that the older standard insecticide formulation, used in a previous study [3], caused adverse effects on lung function and symptoms. The other two standard formulations, which are still commercially available in Australia, had similar, but more severe, adverse effects. The magnitude of the mean change in both lung function and symptoms was small. However, seven (29%) of the subjects exposed to the commercially available aerosols experienced clinically relevant falls in FEV1, suggesting there is potential for a significant clinical problem at high levels of exposure. Unlike the study of Newton and Breslin [3], the present study has shown that exposure to these standard insecticide aerosols was associated with more severe airway responsiveness than was measured after the negative control aerosol. This difference in findings may be due to the fact that in the previous study only four subjects had AHR measured before and after exposure, and the measurement of AHR was made 24 h after the insecticide exposure. In the present study, the effect on AHR was small, approximately half a doubling dose, and was measurable 90 min after completion of the exposure.

The mechanism by which insecticide aerosols affect the airways is unclear. The effect is unlikely to be due to the toxicity of the active ingredients. The dose of insecticide was below the established "No observable effect" level and close to the threshold limit value [4±6]. In insects, pyrethroids kill by a direct effect on the nervous system, but this is avoided in mammals by rapid hydrolysis in the liver [12]. Numerous studies have shown that domestic irritants such as perfumes [13, 14] or hair sprays [15] can cause a change in lung function. None of these studies have examined the effect on AHR, so it is not clear whether these substances can affect AHR directly, and the cause of the change in AHR in the present study remains unclear. Although the insecticide aerosols caused a fall in lung function immediately after the exposure, by the time the challenge test was performed, FEV1 had returned to baseline levels. It is possible that other parameters of lung function remained abnormal at this time, but it seems unlikely that the change in AHR could be explained on airway geometry alone. Pyrethrum is obtained from an extract of the chrysanthemum flower, and has been associated with allergic rhinitis and contact dermatitis [12, 16, 17]. Insecticide aerosols contain purified pyrethrins or synthetic pyrethroids, which are structurally similar to the natural pyrethrin molecule. Despite reports of "allergic" responses to synthetic pyrethroids [12], including a fatality due to sudden bronchospasm [2], no data have been reported to suggest that these responses are due to an immunoglobulin E-mediated allergic response. At present there are insufficient data either from previous studies or the present study to determine the mechanism of the adverse effects of insecticide aerosols on the airways.

The low irritant formulation did not cause significant adverse effects on lung function or symptoms. Furthermore, the data from this sample suggest that, if there is any fall in FEV1 attributable to this test aerosol, it averages <4% in this susceptible population and is not likely to be clinically significant. Comparison of the low irritant formulation with the other sprays shows that it causes a lesser
reduction in lung function than the two commercially-available sprays and less eye symptoms than the older formulation. Other differences between the low irritant formulation and both the older formulation and the commercially-available sprays tended to favour the low irritant formulation but did not reach statistical significance. The active insecticides in the low irritant formulation differed in chemical structure from those in the standard aerosols, but it was beyond the scope of the present study to identify the relevant chemical differences responsible for the different effects.

In conclusion, this study provides objective evidence that subjects who report chest, nose, or eye symptoms following exposure to domestic fly-sprays may be at risk of significant changes in lung function, and an increase in airway responsiveness. The exposure levels used in this study are higher than those which occur with normal domestic use of insecticide aerosols, but could be achieved by over-zealous use in a confined space. Further studies may help to determine whether the mechanism of action is due to an irritant effect of the spray on sensory nerves in the airways, or whether there is any evidence of an immunoglobulin E mediated allergic response. These findings support the view that the low irritant formulation is unlikely to cause clinically significant adverse effects in patients with asthma and that, in this respect, it represents an improvement on the standard formulations.

References