Occupational asthma due to tea dust

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Abstract
Three patients are described who had developed asthma after working as tea packers. In two cases the diagnosis was confirmed by serial monitoring of peak expiratory flow rates, bronchial responsiveness to histamine, and specific inhalation challenges in the laboratory. The third patient experienced isolated changes in bronchial responsiveness to histamine after periods of exposure at work and after specific inhalation challenges in the laboratory without showing spirometric changes. Two of the three subjects were non-atopic; none had an immediate reaction to skin prick testing with a tea solution.

Occupational asthma among workers processing different plants, grains, and beans has been frequently documented.1,2 Occupational asthma due to tea dust, a dust released during the processing of tea leaves, has been described3 and asthma due to tea dust has been documented in a tea packer.4 We describe three patients who developed occupational asthma as a reaction to tea dust while working in a plant where tea dust was packed into tea bags.

Case reports
Baseline data for all the patients are given in the table. Histamine challenge tests were carried out as described elsewhere.2,4

| Subject 1 | This 43 year old woman had been symptom free until nine months before her first visit to the clinic, at which time she had noticed rhinorrhea, cough, dyspnoea, and wheezing. The symptoms were more pronounced at work, and she woke during the night with chest symptoms when she had been at work. Her symptoms improved with oral theophylline, an inhaled beta2 adrenergic agent, and beclomethasone, and they cleared after one month away from work, so that she was able to stop all medication. On her return to work her asthma and rhinitis recurred within two weeks. She was again removed from the workplace for one month, and given a short course of oral prednisone. When she returned to work, taking oral theophylline and inhaled beclomethasone (200 µg daily), symptoms recurred again and were controlled by inhaled salbutamol as needed. She was symptom free when first seen by us, having been away from work for three weeks. Spirometry gave normal results.7 The variability in her peak flow meter values recorded every two waking hours was less than 20% (fig 1). Two histamine inhalation tests showed borderline hyperresponsiveness.

Returning to work caused a recurrence of her symptoms, increased fluctuation in her peak expiratory flow (PEF) and a fall in her PC20 (the provocation concentration of histamine causing a 20%, fall in FEV1) and FEV1, (fig 1). Removal from the workplace caused an improvement in PEF, FEV1, and symptoms without any extra medication; the patient was completely symptom free after six weeks away from work. The FEV1 returned to normal but her PC20 histamine was still low more than a month after she left work.

To determine whether tea dust was responsible for the exacerbation of asthma at work, specific inhalation challenges were performed (fig 2). Exposure to wood dust for two hours,4 as a control, induced no change in spirometric values. PC20 was 1-6 mg/ml at the end of the day. The next day, gradual exposure to tea dust for a total of 30 minutes induced an atypical early late asthmatic reaction, beginning 20 minutes after the last exposure and producing a maximum fall in FEV1 of 29% from baseline three hours after exposure ended. PC20 on the following day was decreased slightly to 0-69 mg/ml (FEV1, within 10% of the baseline value of the first histamine test).

| Subject 2 | This 38 year old woman, a machine operator, reported perennial rhinitis of seven years' duration, associated with cough, shortness of breath, and wheezing over the previous four years. The symptoms increased three hours after arriving at work and caused her to awaken |

Clinical and functional results at the time of first assessment of the three women with occupational asthma

<table>
<thead>
<tr>
<th>Patient No</th>
<th>Age (y)</th>
<th>Atopy</th>
<th>Skinprick reaction to tea dust</th>
<th>Smoking habit</th>
<th>Duration of exposure (y)</th>
<th>Duration of symptoms (y)</th>
<th>FEV1 (% pred)</th>
<th>FEV1/FVC (% pred)</th>
<th>PC20 (mg/ml)</th>
<th>Specific IgE to tea dust (cpm)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>43</td>
<td>-</td>
<td>Ex-smoker</td>
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<td>100</td>
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<td>-</td>
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<td>103</td>
<td>7-4</td>
<td>362</td>
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<tr>
<td>3</td>
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<td>+</td>
<td>Non-smoker</td>
<td></td>
<td>19</td>
<td>&lt;1</td>
<td>95</td>
<td>103</td>
<td>7-4</td>
<td>392</td>
</tr>
</tbody>
</table>

*Mean value of 382 cpm in three non-atopic, non-asthmatic individuals. ND—not done.
occasionally at night. They improved during weekends and holidays. She was treated with oxtriphylline and an inhaled beta$_2$ adrenergic agent. The diagnosis of asthma was confirmed two years after the start of symptoms, when spirometry showed partially reversible airway obstruction with an FEV$_1$ and forced vital capacity (FVC) of 1.35 and 2.02 l, improving to 1.74 and 2.68 l after inhaled beta$_2$ adrenergic agent (predicted values 2.51 and 2.94 l). Her chest symptoms steadily worsened over several years, so she left her job one year before being seen. She was completely symptom free after several months away from work. She underwent specific inhalation challenges as for subject 1. Neither exposure to tea dust for 60 minutes as described above nor inhalation of a tea dust solution for five and 30 minutes via a Wright nebuliser (output 0.14 ml/min) on three consecutive days induced a change in FEV$_1$, of 8% or more over eight hours (fig 3), though she developed cough and shortness of breath. Her PC$_{20}$ fell on each occasion; at this time the maximum variability in baseline FEV$_1$ was 10-9%. The diagnosis was not retained because FEV$_1$ did not change after the challenge test; no explanation was found for the changes in bronchial responsiveness.

Two years later, after a diagnosis of occupational asthma had been confirmed in another worker from the same processing plant (No 3 below), we decided to repeat the tea dust exposure tests at work and in the laboratory for longer periods. The subject was still symptom free and had no bronchial hyperresponsiveness. Returning to work caused her to cough but was not associated with any appreciable changes in peak expiratory flow (PEF). Her PC$_{20}$ fell after two periods of five days at work (fig 3), but had risen to over 32 mg/ml one week later. Specific inhalation challenges were then repeated in the laboratory. Baseline PC$_{20}$ was over 16 mg/ml. Exposure to tea dust for four hours resulted in a fall in PC$_{20}$ from over 32 to 3.8 mg/ml, with progressive recovery over the next six days. The maximum variability in FEV$_1$ during the day was 7-1%. After exposure to wood dust for four hours the maximum variability in FEV$_1$ was 8-6% and PC$_{20}$ was over 16 mg/ml. Repeat exposure to tea dust for four hours did not result in any change in FEV$_1$, (maximum daily variability 4.3%). There was also no change in lung volumes (residual volume, functional residual capacity, total lung capacity) or maximum flow at 50% of forced vital capacity. There was, however, a fall in PC$_{20}$ with progressive recovery over the next few days at a time when the maximum variability in baseline FEV$_1$ before each histamine test was 10-6% (fig 3). The subject had cough and shortness of breath at the time of exposure to tea dust but not with wood dust.

An asthmatic subject with a PC$_{20}$ of 1.4 mg/ml was exposed to tea dust for two hours in the laboratory as a control; the exposure, which was similar to the one for the other subjects, did not induce any changes in FEV$_1$ (<10%) or in PC$_{20}$ (2-1 mg/ml at the end of the day).

FIGURE 1 Combined monitoring of peak expiratory flow (PEF) and responsiveness to histamine (PC$_{20}$) in subject 1. Significant changes in PEF were documented in the second week of exposure, coinciding with a need for an inhaled beta$_2$ adrenergic agent (S). FEV$_1$ values at the time of PC$_{20}$ assessment are given.

FIGURE 2 Results of specific inhalation challenges with wood dust (two hours, ○) and tea dust (30 minutes, ●) in subject 1. S—inhalation of a beta$_2$ adrenergic agent. Specific inhalation challenges were performed during 22 August–2 September.

SUBJECT 3
This 40 year old woman reported cough with chest tightness and wheezing that was more pronounced in the afternoon when she was at work. She also reported waking at night with asthmatic symptoms. There was improvement during weekends and holidays. She was atopic (positive skinprick test responses to ragweed pollen and Dermatophagoides farinae) but reported no history of hay fever. The PC$_{20}$ was 7-4 mg/ml (baseline FEV$_1$, 2.74 l) after two months away from work. After returning to work for two weeks her PC$_{20}$ had fallen to 1.5 mg/ml (baseline FEV$_1$, 2.56 l). PEF monitoring showed greater fluctuations when she was at work than during a weekend and a period before exposure at work (fig 4). Specific inhala-
Occupational asthma due to tea dust were documented. The duration of exposure to each agent at work and in the laboratory is shown.

Discussion

Two cases of occupational asthma due to tea dust have been already documented. Although in the first case the worker was employed in the primary industry, in the second case the worker was a tea packer like our subjects. In the second case the diagnosis was confirmed by serial monitoring of PEF and specific inhalation challenges, whereas in the first case the diagnosis was a clinical one. In our two workers the diagnosis of occupational asthma was confirmed by monitoring peak expiratory flow. As this does not exclude the possibility of an irritant reaction, we combined monitoring with serial assessment of bronchial responsiveness. Substantial changes in PC_{20} were documented in both subjects and were prolonged in the first subject. Finally, specific inhalation challenges confirmed the diagnosis of occupational asthma as they induced a late reaction in one subject and an atypical immediate or early late reaction in the other. Although the exposure level at the time of specific inhalation challenges was not monitored and could have been high at this time, this pattern of reaction excludes a non-specific irritant mechanism; asthmatic subjects exposed to high levels of particles such as sawdust do not generally show changes in spirometric values and bronchial responsiveness after exposure. Furthermore, a control asthmatic subject showed no changes in FEV_{1} or PC_{20} after a similar exposure to tea.

The second subject illustrates an interesting point. Although exposure to tea dust caused little change in FEV_{1} or PEF, there was a change in PC_{20} of up to four doubling doses, from normal to within the asthmatic range. Such changes cannot be attributed to an irritant reaction; the subject did not show bronchial hyperresponsiveness at the start of the challenges on three separate occasions and recovery of PC_{20} took several days. Finally, changes in PC_{20} were not documented in a control subject who had bronchial hyperresponsiveness. The effect of exposure to an environmental asthma inducing agent may at times be detected more readily from change in bronchial responsiveness than from changes in FEV_{1} or FVC. The lack of changes in FEV_{1} and/or FVC after exposure to tea dust might be due to the fact that the subject had been away from work for a long interval when the tests were carried out. She is likely to have lost some sensitisation. It is difficult to label this case occupational asthma, though she

![Figure 3](image-url) Results of specific inhalation challenges in subject 2. No significant changes in FEV_{1} were documented. The duration of exposure to each agent at work and in the laboratory is shown.

![Figure 4](image-url) Monitoring of peak expiratory flow (PEF) at work and away from work in subject 3. S—inhalation of a beta_2-adrenergic agent.

![Figure 5](image-url) Results of specific inhalation challenges with wood dust (O) and tea dust ( ) first exposure, ( ) second exposure) in subject 3. S—inhalation of a beta_2-adrenergic agent.
definitely had asthma when she was working at the tea plant, as documented by her reversible airway obstruction. There is a similar case of a snow crab worker in whom the first specific challenge, done after several months away from work, gave negative results but who developed asthma again several weeks after returning to work; challenges at that time gave positive results. This has also been reported in a worker exposed to isocyanate. For us to prove the point definitively our subject would have had to return to work for several weeks or months.

That two of the three workers started having symptoms after stopping smoking is interesting. Smoking is inconsistently related to occupational asthma, but asthma that occurs or recurs after cessation of smoking has been described; the mechanism is unknown.

The mechanism of this type of occupational asthma is not understood. We and others were unable to detect specific IgE, and skin tests failed to elicit an immediate reaction. The causative agent could be the tea plant itself or a microbial contaminant, though immediate skin reactions were found in the patient in the original report. The prevalence of occupational asthma among workers exposed to tea dust remains to be explored.

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