Seasonal variation in non-specific bronchial reactivity: a study of wheat workers with a history of wheat associated asthma

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ABSTRACT To investigate seasonal variation in non-specific bronchial reactivity in wheat workers, we carried out histamine inhalation tests in 29 workers (28 of them men) from a small farming community with symptoms of wheat associated asthma before, during and after the 1983–4 Australian wheat harvest season. Four were cigarette smokers, and the age range was 12–54 (mean (SD) 30 (10)) years. Twenty-eight subjects were atopic (one positive skin prick test result in tests with 10 common antigens), 60% reacting to house dust mite and all to at least one of eight wheat antigens. Baseline spirometry gave normal results (mean FVC, 90% (SD 8%) predicted; FVC 91% (7%) predicted). Bronchial reactivity was tested by the method of Yan et al. The cumulative doses of histamine acid phosphate (up to 3·91 μmol) that caused a fall of 20% from baseline in FEV₁ was determined (PD₂₀) and expressed as the geometric mean. In the low exposure season, May 1983, nine subjects had a PD₂₀ (mean 1·2, range 0·3–3·9 μmol). The number rose to 19 in the summer harvest season, December 1983 (mean 0·8, range 0·07–3·9 μmol) and returned to nine in the subsequent winter, July 1984 (mean 1·8, range 0·4–3·9 μmol). The change in the number of subjects with a PD₂₀ was significant (p < 0·01). Four additional subjects probably had increased bronchial reactivity in the harvest season: in two the post-saline FEV₁ was too unstable to give them histamine challenge and in two the challenge was inadvertently discontinued prematurely. Baseline FEV₁ and FVC fell by 8% between the first and second studies (p < 0·001); values were intermediate in the third study (FEV₁, 3·74, 3·44, and 3·57; FVC 4·66, 4·28, and 4·41 litres respectively). Linear modelling analysis of log PD₂₀, season, FEV₁, FVC, age, seasonality of asthma symptoms and skin test data indicated that the harvest season was the only significant determinant of variation in log PD₂₀. It is concluded that in these wheat workers there is a seasonal variation in bronchial reactivity that may reflect a response to allergens associated with grain.

Asthma is acknowledged to be a considerable health problem in an increasing number of occupations.1,2 The harvesting, drying, and transportation of grain, its milling, and the handling of flour have all been associated with illness of both the upper and the lower respiratory tracts.3–6

An increase in non-specific bronchial reactivity is generally accepted as present in most if not all patients with current or symptomatic asthma7 and it has been suggested that the measurement of non-specific bronchial reactivity should be an essential requirement for establishing the diagnosis of asthma, especially in epidemiological studies.8 Non-specific bronchial reactivity is not stable, however, and it has been shown to vary spontaneously, with respiratory infections, with allergen exposure, with treatment, and with the severity of asthma as reflected by drug requirements or symptoms.9–15

In this study 29 wheat workers with a history of asthma like symptoms on exposure to wheat were
investigated for changes in non-specific bronchial reactivity before, during, and after the Australian wheat harvest season of 1983-4. Our major hypothesis was that seasonal exposure to wheat would be associated with a seasonal change in bronchial reactivity.

Methods

The study was carried out in the wheat district of New South Wales in a town of about 1900 people situated 450 kilometres north west of Newcastle. The 29 subjects were studied on each of three occasions: May 1983 (late autumn), December 1983 (summer), and July 1984 (winter).

In May 1983 we studied 66 local inhabitants who had either upper or lower respiratory tract symptoms on exposure to wheat. Of these, 57 had symptoms consistent with asthma according to the criteria of the American Thoracic Society.16 Of the subjects with asthma, 33 volunteered to be studied in December 1983 and 31 in July 1984. We report the results on 29 subjects who participated on all three occasions. All subjects gave informed consent. The project was approved by the ethics committee of the University of Newcastle.

Symptoms and Questionnaire

Asthma was classified according to whether it occurred only on exposure to grain or at other times throughout the year, such as with a respiratory tract infection. This latter group was designated “Perennial.” The timing of symptoms of cough, wheeze, and dyspnoea in relation to exposure to grain was classified as “immediate” if they occurred on contact or “delayed” if symptoms came on later that day or evening. Each subject completed the ATS-DLD 78 questionnaire during the first study, though none of these data have been used in this analysis. Since we could not validly document asthma symptoms during the entire harvest season (December 1983–January 1984), we did not try to use asthma like symptoms for any purpose other than to select subjects.

Spirometry

Forced vital capacity (FVC) and forced expiratory volume in one second (FEV1) were measured with a dry wedge spirometer (Vitalograph) calibrated with a three litre syringe. Volumes were not corrected to BTPS because of the evidence that expired air does not cool to ambient temperature with a Vitalograph and that full correction to BTPS gives incorrectly large values.17 The best FEV1 and FVC of at least three manoeuvres were recorded as the baseline values on each occasion. Predicted values were calculated from the formulae of Knudson et al.18

Bronchial Reactivity

Non-specific bronchial reactivity was measured by a histamine inhalation test with the method of Yan et al.19 Following a control inhalation of nebulised saline, histamine acid phosphate was administered by a cumulative dose technique from four DeVilbiss nebulisers containing concentrations of 3-13, 6-25, 25, and 50 mg/ml histamine acid phosphate. The procedure was stopped when the FEV1 had fallen by 20% or more from the baseline. All subjects whose FEV1 did not fall by 20% received a cumulative dose of 3-91 μmol of histamine. Nebulised salbutamol was administered after the histamine inhalation test to all subjects whose FEV1 had fallen by 10% or more. Bronchial reactivity was calculated from a plot of change in FEV1 from baseline against log-dose of histamine. The dose at which FEV1 was 20% lower than baseline (PD20) was interpolated from the graph and expressed as the geometric mean.

Skin Prick Tests

During the third visit skin prick tests were carried out with 10 common antigens (Dermatophagoides pteronyssinus, D farinae, Aspergillus sp, mould mix, perennial rye grass, Timothy, seven grass mix, plain tain, dog, cat—Hollister-Stier). Atopy was defined as the presence of at least one positive skin test response (weal diameter of 3 mm or more at 15 minutes). Skin prick tests were also carried out with eight wheat antigens including commercially available extracts of cultivated wheat and wheat smut (Hollister-Stier) as well as extracts prepared in our own laboratory—wheat whole grain, wheat chaff, wheat straw dust—wheat grain dust, wheat header dust, wheat silo dust—according to the method of Baldo and Wrigley.19

Statistical Analysis

For comparison of grouped data for seasonal change in bronchial reactivity, a Pearson χ2 test was performed. Comparison of changes in spirometric values were carried out by analysis of variance and paired t test. To explore the determinants of the changes in PD20 a regression analysis was carried out with the linear modelling program known as GLIM.20 The factors entered into the analysis included the results on the three occasions for FEV1, FVC, and log PD20 as well as age, asthma classified as “perennial” or “grain only” and a positive skin prick test reaction to house dust mite.
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Results

DEMOGRAPHIC DETAILS AND BASELINE SPIROMETRY
Of the twenty-nine subjects, there were nineteen farmers, three farm equipment mechanics, two ex-farmers, two students, one farm-hand, one ex-wheat-grader and one stock carrier. There was one woman and the ages ranged from 12 to 54 (mean 30) years. There were four cigarette smokers. Eighteen subjects had symptoms on exposure to grain but not with any other precipitant (classified as "grain only"). Thirteen subjects (seven with "grain only" and six with "perennial" asthma) had only immediate symptoms on exposure to grain.

TREATMENT for asthma symptoms within the month preceding each study was as follows. May 1983 (autumn): 22 no treatment; 4 bronchodilator; 3 bronchodilator plus sodium cromoglicate or corticosteroids. December 1983 (summer harvest): 13 no treatment; 11 bronchodilator; 5 bronchodilator plus sodium cromoglicate or corticosteroids. July 1984 (winter): 12 no treatment; 15 bronchodilator; 2 bronchodilator plus sodium cromoglicate or corticosteroids. No subjects took a bronchodilator within six hours of the histamine challenge. Further details are available on request.

Baseline spirometric values in the first study were within normal limits (greater than 80% of the predicted value for FEV₁ and FVC) for 27 of the 29 subjects. A 25 year old man had an FEV₁ of 77% predicted and a 44 year old woman had an FEV₁ of 70% predicted. Neither smoked. For the 29 subjects the mean FEV₁ as a percentage of the predicted value was 90.4% (SD 8.0%) and the mean FVC 91.3% (7.2%).

At the second study, during the harvest season, there was a fall of 8% in mean FEV₁ (from 3.74 (SD0.6) to 3.44 (0.6) litres; p < 0.001), with a subsequent rise at the third study (3.57 (0.59) litres: p = 0.04). Changes in FVC were similar to those seen with FEV₁ (from 4.66 (0.73) to 4.28 (0.63) to 4.41 (0.68) l). The data for FEV₁ and FVC on the third occasion were from 28 subjects because one subject was unable to produce a repeatable forced expirogram.

SKINPRICK TESTS
Twenty-eight of the eighty-nine subjects had at least one positive skin test reaction to common allergens, the number of positive values ranging from 1 to 9 (mean 5 (SD 2.5)). Fifty nine per cent of subjects had a positive reaction to house dust mite. Comparison of the skin test results of the "grain only" and the "perennial" groups showed no substantial differences (for instance, "grain only" 61% positive for house dust mite, total positive results 4 (SD 2); “perennial” 55% positive for house dust mite; total positive results 5 (3)). All subjects had at least one positive reaction to one of eight wheat antigens. The prevalence of positive results ranged from 48% for wheat smut to 86% for wheat straw dust.

HISTAMINE INHALATION TEST
There was a change in bronchial reactivity between the harvest and non-harvest seasons when the results were analysed for the number of subjects with a PD₂₀ (χ² = 6.91, p < 0.01; see table). Nine subjects had a PD₂₀ on the first occasion (geometric mean 1.2, range 0-3-3-9 μmol/l), 19 on the second (mean 0-8, range 0-07–3-9 μmol/l), and nine on the third (mean 1-8, range 0-4–3-9 μmol/l).

On the second occasion (during the harvest season) there were two subjects whose post-saline FEV₁ was too unstable for the histamine inhalation test and two whose test was inadvertently stopped before a 20% fall in FEV₁ (FEV₁ had fallen by 10% or more after only 0-06 and 0-62 μmol of histamine). On the third occasion (winter 1984) one subject’s baseline spirometric values were too unstable for the histamine inhalation test.

The figure shows results for PD₂₀ for 21 subjects. Omitted are five whose results were negative on all occasions and the three subjects with missing data described above. Two of the three subjects in the figure whose results were negative in the harvest season may have had a measurable PD₂₀ if the histamine inhalation test had not been stopped prematurely.

ANALYSIS OF DETERMINANTS OF BRONCHIAL REACTIVITY
When a linear model of the data was constructed for age, type of asthma, presence or absence of atopy, and the results of PD₂₀, FEV₁, and FVC for the different seasons, the only significant explanatory variable for the variation in log PD₂₀ was season—that is, the second occasion, during the harvest season, was a significant determinant (t = 3.86, p < 0.001).

Spirometric and bronchial reactivity data

<table>
<thead>
<tr>
<th></th>
<th>May 1983</th>
<th>Dec 1983</th>
<th>July 1984</th>
</tr>
</thead>
<tbody>
<tr>
<td>FEV₁ (l, mean (SD))</td>
<td>3.74 (0.61)</td>
<td>3.44 (0.61)</td>
<td>3.57 (0.59) *</td>
</tr>
<tr>
<td>FVC (l, mean (SD))</td>
<td>4.66 (0.73)</td>
<td>4.28 (0.62)</td>
<td>4.41 (0.68) *</td>
</tr>
<tr>
<td>No with positive HIT</td>
<td>9</td>
<td>19</td>
<td>9</td>
</tr>
<tr>
<td>PD₂₀ (μmol, geometric mean (range))</td>
<td>1.2 (0-3-3-9)</td>
<td>0.8 (0-07–3-9)</td>
<td>1.8 (0-4–3-9)</td>
</tr>
</tbody>
</table>

*28 subjects only.

FEV₁, forced expiratory volume in one second; FVC, forced vital capacity; positive HIT, positive result in histamine inhalation test—PD₂₀ < 3-9 μmol histamine; PD₂₀ dose of histamine causing a fall in FEV₁ of 20%. 

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Discussion

We have shown that in the decreased and the bronchial ced methodological certain our sample our season. cannot validly the with bronchial limitations, symptoms. We did bronchial reactivity study during toms. These include of the previous results. These include the representativeness of the sample of subjects, the validity of the method for measuring bronchial reactivity, and the possible effect of reduction in lung function during the harvest season.

Since our subjects were all volunteers who had symptoms suggestive of asthma in the two years preceding our first test, they may not be representative of farmers in either the community or the state. These results cannot therefore be generalised to the wheat farming community. Furthermore, in the absence of measurements in a symptomless group we cannot validly assume any causal relationship between the increase in bronchial reactivity and the asthma like symptoms reported for the previous seasons. Despite these limitations, the results support the hypothesis that bronchial reactivity in wheat workers does change with the seasons coincidentally with changes in their symptoms. We did not ask about respiratory symptoms during the 1983–4 harvest season since the bronchial reactivity study was carried out early in the season and subjects who had symptoms later in the season would have been misclassified.

The technique for measurement of bronchial reactivity used in the present study has been extensively evaluated by Yan and colleagues and more recently by Britton et al. We have carried out our own quality control studies, which showed reproducibility to be within one doubling dose of histamine in a group of volunteers from the medical school. We agree with the conclusion of Britton et al. that the Yan technique is a good test for use in epidemiological studies of bronchial reactivity.

An important consideration in the interpretation of our data is whether the 8% fall in FEV, found in the second test could have affected the measurement of bronchial reactivity, via a change in airway geometry. To our knowledge there is no way to resolve this question in quantitative terms, but such a relatively small fall in FEV, seems unlikely to be associated with a substantial enough change in geometry to cause the variation in bronchial reactivity that we have measured. In support of this argument are the results from the third occasion, when there was a reversal of the increase in bronchial reactivity without complete return of lung function to previous values, the FEV, being on average 4.5% below that of the first occasion. Furthermore, the linear model analysis showed a highly significant effect of the season but not FEV, on the variation of PD20.

We conclude that the change in bronchial reactivity that we observed is real and not explained by any artefact. What is the possible mechanism of the change? There is a reasonable amount of evidence to support the hypothesis that an allergen induced inflammatory process may be responsible. Such a mechanism has been proposed by Boulet et al. and by Sotomayor et al. for the seasonal changes they found in bronchial reactivity in patients with allergy to grass pollens. The case for the role of inflammation was strengthened by the finding in the latter study that the changes in bronchial reactivity were reversed by methylprednisolone 16 mg/day but not by placebo. Since all but one of our subjects reacted to common antigens and all to wheat antigens, we believe that a similar mechanism may be operating in our subjects.

While the major aim of our study was to describe the seasonal variation in bronchial reactivity in wheat workers, our results can be used to assess the predictive value of bronchial reactivity tests done before exposure to an environment which is suspected to provoke asthma. Although all subjects had a history of asthma like symptoms in the previous two seasons, only nine out of 29 subjects had a measurable PD20 in the autumn before the harvest season. Over half of the “negative” subjects subsequently developed a measureable PD20 in the harvest season. These results,
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admittedly from a small sample, indicate that bronchial reactivity is not a biologically stable entity and bring into question its use except for research. There is still uncertainty about the appropriate role of bronchial reactivity in occupational medicine; the evidence is well summarised in a recent review.2

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