Two variants of occupational asthma separable by exhaled breath nitric oxide level

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ABSTRACT

Exhaled nitric oxide (F\textsubscript{ENO}) has been used as a marker of asthmatic inflammation in non-occupational asthma, but some asthmatics have a normal F\textsubscript{ENO}. In this study we investigated whether, normal F\textsubscript{ENO} variants have less reactivity in methacholine challenge and smaller peak expiratory flow (PEF) responses than high F\textsubscript{ENO} variants in a group of occupational asthmatics.

Methods: We measured F\textsubscript{ENO} and PD\textsubscript{20} in methacholine challenge in 60 workers currently exposed to occupational agents, who were referred consecutively to a specialist occupational lung disease clinic and whose serial PEF records confirmed occupational asthma. Bronchial responsiveness (PD\textsubscript{20} in methacholine challenge) and the degree of PEF change to occupational exposures, (measured by calculating diurnal variation and the area between curves score of the serial PEF record in Oasys), were compared between those with normal and raised F\textsubscript{ENO}. Potential confounding factors such as smoking, atopy and inhaled corticosteroid use, were adjusted for.

Results: There was a significant correlation between F\textsubscript{ENO} and bronchial hyperresponsiveness in methacholine challenge (p=0.011), after controlling for confounders. Reactivity to methacholine was significantly lower in the normal F\textsubscript{ENO} group compared to the raised F\textsubscript{ENO} group (p=0.035). The two F\textsubscript{ENO} variants did not differ significantly according to the causal agent, the magnitude of the response in PEF to the asthmagen at work, or diurnal variation.
Conclusions: Occupational asthma patients present as two different variants based on 
FE\textsubscript{NO}. The group with normal FE\textsubscript{NO} have less reactivity in methacholine challenge, while 
the PEF changes in relation to work are similar.
INTRODUCTION

Measurement of exhaled breath nitric oxide (FE\textsubscript{NO}) has been promoted as a measure of airway inflammation in asthma [1-4]. It has been shown to be correlated with sputum eosinophilia and non specific reactivity in asthmatics [5-14] but has the advantages of being less invasive for the patient and less labour intensive for the clinician. However, some symptomatic asthmatics have been reported to have normal levels of FE\textsubscript{NO} [9-15-17] even when factors such as inhaled corticosteroid therapy and smoking have been accounted for. In the diagnosis of occupational asthma, one of the best first line investigations for occupational asthma is serial peak expiratory flow (PEF) monitoring and is recommended by several guidelines [18;19]. It has been suggested previously that using changes in sputum eosinophil counts between periods of exposure and non-exposure increases the sensitivity and specificity of serial PEF measurement in the diagnosis of occupational asthma [20]. Specific inhalation challenge tests to occupational agents have resulted in a mean increase of exhaled nitric oxide levels [21-24]. However, some workers with positive challenges have not showed changes. We have previously found a strong positive correlation between exhaled nitric oxide level and sputum eosinophil count in workers with occupational asthma exposed to low molecular weight agents and a relationship between sputum eosinophilia and non specific reactivity [9]. The study suggested that workers can be separated into two variants, those with eosinophilic Airways inflammation and those with non-eosinophilic inflammation and that they would also be separable by FE\textsubscript{NO} due to the strong relationship between the two indices. The aim of this study was to see whether our retrospective analysis could be confirmed with a prospective group, and whether and whether the magnitude of PEF response to occupational exposure is related to FE\textsubscript{NO}. 
METHODS

Study Population

Consecutive workers referred to the Occupational Lung Disease Clinic, Birmingham, UK between November 2001 and December 2004 were recruited who had performed an exhaled nitric oxide measurement (FeNO), methacholine challenge test and serial PEF record while still exposed at work. Sixty subjects whose serial PEF measurements showed occupational asthma while exposed to the causative agent and who had a diagnosis of occupational asthma formed the study population. The study was approved by the East Birmingham Local Ethics Committee (reference 929).

Measurements

Workers were requested to record PEF every 2 hours from waking to going to bed on work days and days away from work for a total of 4 weeks. The best of 3 PEF readings were recorded on each occasion, provided that the best 2 readings were within 20 l/min of each other. Records were plotted, linearised [25] (if recorded on a non-linear PEF meter) and analysed by the Oasys computer program [26]. Those with a work effect index score $\geq 2.51$, (that was used as a cut-off point for definite occupational effect) [26] were included in this analysis.

Spirometry, FeNO and non-specific bronchial reactivity in methacholine challenge were performed within 24 hours of work exposure after withholding treatment with long acting $\beta$-agonists for 24 hours (including combined steroid and long acting $\beta$-agonists inhalers), short-acting $\beta$-agonists for 6 hours and tiotropium for 36 hours as part of their routine clinic visit.
Spirometry was performed on either a wedge bellows Vitalograph spirometer or on the Jaeger pulmonary function system according to ERS/ATS standards [27]. Non-specific bronchial reactivity to methacholine was measured using the Yan technique [28]. \( \text{FE}_{\text{NO}} \) was measured during exhalation at 50ml/second using the Niox from Aerocrine, which requires values from two readings to be within 10% as recommended by the ATS/ERS [29] and performed before spirometry. The Oasys program [26] was used to calculate diurnal variation on days at and away from work and the area between curves (ABC) based on mean PEF on work days and days away from work (ABC score) plotted by waking time (Figure 1) [30].

**Figure 1.** The ABC plot of a worker exposed to chrome from stainless steel welding.

He has normal methacholine reactivity (>4800mcg) and an \( \text{FE}_{\text{NO}} \) of 6.1ppb. The plot has a 56 L/min/hour difference between the mean curves of PEF on work and rest days. In the bottom panel, the first row of numbers is the time from waking in 2-hourly sections e.g. 00-02; 02-04 etc. The second row shows the number of readings used for the mean PEF curves in each 2-hourly section (left side shows work readings and right side shows rest readings). The third row shows the area between the curves for each 2-hourly section which are then used to calculate the ABC score which is in litres/min/hour. A score of \( \geq 15 \) L/min/hr has a sensitivity of 69% and specificity of 100% for occupational asthma diagnosis [30].
Workers were split into normal and raised nitric oxide level groups based on an eosinophil cut off of 2.2% which was used in our previous study to separate eosinophilic and non eosinophilic variants [9]. A cut off of 14.7ppb for smokers and 22.1ppb for non smokers (equivalent to \(<\) or \(\geq\) 2.2% sputum eosinophilia) was selected from a regression analysis of all our previous combined measurements of sputum eosinophils and F\(_{\text{ENO}}\). These values were then used to separate workers into those with normal F\(_{\text{ENO}}\) and those with raised F\(_{\text{ENO}}\) levels.

Characteristics of the workers such as smoking history, atopy (defined as at least one positive skin prick test of \(\geq 3\)mm wheal to a common environmental allergen using saline and histamine as negative and positive controls) and inhaled corticosteroid treatment were recorded. Inhaled corticosteroids were classified into groups according to the GINA guidelines [31] for analysis against F\(_{\text{ENO}}\).

**Statistical analysis**

Data was analysed by using F\(_{\text{ENO}}\) as a continuous variable and also by grouping the workers into two variants based on their F\(_{\text{ENO}}\) level. Physiological data were not normally distributed, so reactivity to methacholine and nitric oxide levels were log transformed. Subjects who had a PD\(_{20}\) >4,800 \(\mu\)g (the highest dose used) in methacholine challenge had their percent fall in FEV\(_1\) extrapolated to give a PD\(_{20}\) value. Differences in physiological parameters between groups were assessed using a Mann Whitney U test or Chi-square test for non-parametric data and either independent t-test or one way ANOVA for parametric data (age, FEV\(_1\) percent predicted, ABC PEF score and log transformed reactivity to methacholine and nitric oxide). Multiple linear regression was used for controlling for variables potentially confounding the relation between F\(_{\text{ENO}}\) and bronchial
hyperresponsiveness. Pearson correlation was used to compare reactivity to methacholine and nitric oxide levels when using both as continuous data. The Yates’ continuity correction was used when at least one cell count was <5 when performing the Chi-square statistic. SPSS version 15 was used for all statistics.

RESULTS
Workers had a mean age of 44 years and 83% were males. Mean FENO levels were similar between atopics and non-atopics (p=0.521), males and females (p=0.183) and those with an FEV1 percent predicted of <80% or >80% (p=0.547). There were eighteen workers at Step 4 of the GINA treatment pathway, eleven at step 3, eight at step 2 and twenty-three on inhaled short acting beta agonists only. There was no difference in log FENO between these groups (p=0.591). Current smokers had significantly lower nitric oxide levels (p=0.013) compared to ex or never smokers. Those who showed bronchial hyperresponsiveness in methacholine challenge had a significantly higher FENO (p=0.006). Table 1 shows statistical comparisons of characteristics and physiological parameters between raised and normal FENO groups, using the different cut-off points for smokers and ex or never smokers.
Table 1. Characteristics of the two variants of occupational asthma separated by FE\textsubscript{NO} level and smoking

<table>
<thead>
<tr>
<th></th>
<th>Normal FE\textsubscript{NO} (smokers &lt;14.7ppb; never/ex &lt;22.1ppb) n=25</th>
<th>Raised FE\textsubscript{NO} (smokers ≥14.7ppb; never/ex ≥22.1ppb) n=35</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean Age (SD)</td>
<td>43.4 (9.7)</td>
<td>45.3 (10.1)</td>
<td>0.469</td>
</tr>
<tr>
<td>% Male</td>
<td>84.0</td>
<td>82.9</td>
<td>0.907</td>
</tr>
<tr>
<td>Mean FEV1 % predicted (SD)</td>
<td>90.7 (21.8)</td>
<td>88.5 (18.1)</td>
<td>0.665</td>
</tr>
<tr>
<td>% Atopic</td>
<td>56.0</td>
<td>62.9</td>
<td>0.593</td>
</tr>
<tr>
<td>% using ICS</td>
<td>54.2</td>
<td>65.7</td>
<td>0.372</td>
</tr>
<tr>
<td>Mean ABC PEF Score (SD)</td>
<td>38.5 (23.9)</td>
<td>29.6 (24.9)</td>
<td>0.196</td>
</tr>
<tr>
<td>Mean PEF work diurnal variation (SD)</td>
<td>17.8 (9.4)</td>
<td>20.5 (12.4)</td>
<td>0.653</td>
</tr>
<tr>
<td>Mean PD20 in Methacholine challenge µg (SD)</td>
<td>5730 (4975)</td>
<td>3883 (5048)</td>
<td>0.035</td>
</tr>
</tbody>
</table>

There was a significant positive correlation between reactivity to methacholine and nitric oxide level when both were analysed as continuous data (Pearson correlation =-0.320; p=0.013). When controlling for smoking, inhaled corticosteroid use and atopy (the main determinants of nitric oxide levels) in multiple linear regression, there was still a significant relationship (R\textsuperscript{2}=0.221; p=0.009). Figure 2 shows the relationships split by current smokers and ex/ never smokers.
Correlations between nitric oxide level and ABC score (as a measure of PEF response) were analysed using multiple linear regression controlling for smoking, inhaled corticosteroid use and atopy. There was not a significant relationship (p=0.781). The ABC score was also compared between those with raised and normal $\text{FE}_{\text{NO}}$ levels in a group of non-smokers who were not taking inhaled corticosteroids. The ABC score was similar (p=0.912). Diurnal variation in PEF was also similar between the two groups (p=0.653).
Workers were analysed for differences in the raised and normal nitric oxide groups according to causative agents (Table 2). There were no differences between those with raised and normal FeNO for high versus low molecular weight agents (P=0.898).

Table 2. Causative occupational exposures by normal and raised FeNO levels.

<table>
<thead>
<tr>
<th>Type of occupational exposure</th>
<th>Normal FeNO (smokers &lt;14.7ppb; never/ex &lt;22.1ppb) n=25</th>
<th>Raised FeNO (smokers ≥14.7ppb; never/ex ≥22.1ppb) n=35</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Metals</td>
<td>9</td>
<td>11</td>
<td>0.711</td>
</tr>
<tr>
<td>Biocides</td>
<td>3</td>
<td>6</td>
<td>0.855</td>
</tr>
<tr>
<td>Metal-working fluid</td>
<td>1</td>
<td>5</td>
<td>0.383</td>
</tr>
<tr>
<td>Isocyanates</td>
<td>3</td>
<td>6</td>
<td>0.855</td>
</tr>
<tr>
<td>Adhesives</td>
<td>2</td>
<td>2</td>
<td>1.000</td>
</tr>
<tr>
<td>Plastics</td>
<td>0</td>
<td>1</td>
<td>1.000</td>
</tr>
<tr>
<td>Other low molecular weight agents</td>
<td>3</td>
<td>0</td>
<td>0.133</td>
</tr>
<tr>
<td>High molecular weight agents</td>
<td>4</td>
<td>4</td>
<td>0.898</td>
</tr>
<tr>
<td>Low molecular weight agents</td>
<td>21</td>
<td>31</td>
<td></td>
</tr>
</tbody>
</table>

**DISCUSSION**

In our study of 60 patients with occupational asthma confirmed by their PEF record, we found that occupational asthma patients can be divided into two variants by FeNO level and that the group with raised FeNO has significantly more reactivity in methacholine
challenge. The two variants do not differ significantly according atopy, causative agents of occupational asthma, inhaled steroid use, or FEV₁ percent predicted, indicating that these did not explain the relation between FEENO and bronchial hyperresponsiveness. Both FEENO groups were similar with respect to changes in PEF in response to occupational exposure, as small and large changes in mean PEF and low and high diurnal variation were seen equally in both normal and raised FEENO groups.

Our results are compatible with others but our interpretation differs. Several groups have shown that the mean FEENO increases with exposure in occupational settings, and that there is a relationship between FEENO and non-specific bronchial reactivity in occupational and non-occupational groups [21;32-39]. Barbinova and Baur found that 52% of occupational asthmatics who had non-specific bronchial hyperresponsiveness had a >50% increase in FEENO post specific inhalation challenge test compared to 20% with normal hyper-responsiveness [40]. The mean changes have however been driven by a subset who show changes, the subgroup without changes in FEENO have not been analysed separately by others.

This study was designed as a follow on to the original Anees et al study [9]. The original observation was from a retrospective analysis, whereas the current paper is wholly prospective data. We started with the hypothesis generated by our previous study that there were two variants of occupational asthma separated by FEENO values that were raised or within normal ranges while exposed, and hypothesised that the response to occupational exposures might differ. By analysing this prospective group, we have confirmed that the two variants differ in non-specific bronchial reactivity, but have not found differences in either the agents responsible for the occupational asthma nor the
responses seen in the workplace measured through serial PEFs. The results indicate that \( \text{FeNO} \) can be used without the need to measure sputum eosinophilia, the former being a simple and cost effective clinical measurement and the latter a much more time-intensive process. There are centres around the world who believe that increased non-specific bronchial reactivity is essential for the diagnosis of occupational asthma. In our experience, normal non-specific reactivity is found in \( \sim 30\% \) of workers currently exposed who have occupational asthma. The results therefore support the inclusion of workers with normal non-specific bronchial reactivity within the family of occupational asthma due to sensitisation. Our PEF response results agree with other studies that have also not shown any correlation between \( \text{FeNO} \) and the magnitude of lung function (mainly \( \text{FEV}_1 \)) in non-occupational asthma [41-43].

We think that these two variants of occupational asthma separable by the \( \text{FeNO} \) level may be related to different types of inflammation in the airways, the raised \( \text{FeNO} \) being related to eosinophilic inflammation and the normal \( \text{FeNO} \) perhaps to neutrophilic or other types of inflammation, which has also been proposed by Taylor et al [44]. This hypothesis is supported by our previous finding that raised \( \text{FeNO} \) was significantly correlated with sputum eosinophilia [9]. Others have also found a linkage between eosinophilia and raised nitric oxide levels [45-47]. We originally hypothesises that the occupational asthmatics with large changes in PEF related to work exposure were more likely to have a raised FeNO than the group with small changes; this however was not supported by our data. Whether these two variants of occupational asthma according to \( \text{FeNO} \) level have implications for prognosis or treatment of the disease needs to be addressed in future studies. One of the factors relating to prognosis (FEV1% predicted) showed similar means for those with raised and normal \( \text{FeNO} \) levels indicating that prognostic factors
may only explain a small amount of the differences in the two variants. This outcome was significantly different in the original retrospective cohort, but other prognostic factors (length of symptomatic exposure and time from first exposure to disease onset) were similar between eosinophilic and non-eosinophilic groups.

A number of studies have shown that inhaled corticosteroid use results in a fall in FE_{NO} levels in patients with asthma [48-55]. As the group with a raised FE_{NO} were on more inhaled ICS than the normal group, we were unable to find a correlation between ICS use and FE_{NO}. A small number of patients may have been misclassified in the normal FE_{NO} group because of this. Workers taking combination inhalers (steroid and long acting beta agonists) would have withheld therapy for 36 hours prior to the clinic appointment for uncompromised non-specific reactivity measurements which may have led to higher FE_{NO} levels in this group. We also found that atopics and non atopics had similar FE_{NO} levels whereas other groups studying asthmatics have found a difference [56-60]. This may be due to the fact that our cohort is a group of occupational asthmatics which may be acting differently to general asthmatics.

**Validity Issues**

All workers in our study had PEF records showing a work-rest pattern compatible with occupational asthma and Oasys score >2.51 (sensitivity of 76% and specificity of 94% for occupational asthma [26]). Workers were recruited consecutively and were currently exposed to the suspected occupational agent at the time of all investigations. There were 12 workers with normal FE_{NO} levels who had a normal reactivity to methacholine and an FEV_{1} percent predicted >80%. Although some may regard these subjects as not having occupational asthma, all of them did fulfil the usual definitions of asthma requiring
airflow obstruction which varies over short periods of time (here within 24 hours of occupational exposures) and their mean diurnal variation at work was 15%. All workers also had a clear, relevant symptom history compatible with occupational asthma and many were exposed to well known causative agents. In addition, 3 had positive specific inhalation challenge tests to the relevant occupational allergen.

Using a cut off for FE\textsubscript{NO} may have it’s limitations, however we believe that by choosing a previously validated cut off based on sputum eosinophilia, this problem has been addressed. With a sample size increase, we may have seen more difference between groups, although looking at the data we feel this is unlikely.

CONCLUSIONS

We have identified two variants of occupational asthma which cannot be separated according to the degree of asthmatic reaction induced by workplace exposures or the agents that they are exposed to, but can be separated by measurement of exhaled nitric oxide whilst symptomatic. The group with raised FE\textsubscript{NO} levels have greater reactivity to methacholine compared to those with normal FE\textsubscript{NO}. This could reflect different types of airway inflammation in these two groups. Whether they differ in prognosis remains a question to be addressed in future studies.

ACKNOWLEDGMENTS

Vicky Moore is the principal data collector, data analyser and principal author for this study. Wasif Anees proposed the original concept, contributed to the collection of data, revised the paper and approved the final version. Professor Burge and Maritta Jaakkola have contributed to the design of the study, interpreted data, revised the paper and
approved the final version. Cedd Burge has contributed to the analysis of data, revised the paper and approved the final version. Alastair Robertson has contributed to the collection of data, revised the paper and approved the final version.

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COMPETING INTERESTS

Conflicts of Interest: Authors Vicky Moore, Maritta Jaakkola, Cedd Burge, Wasif Anees and Alastair Robertson have no conflicts to disclose for this manuscript. Professor Burge promotes and disseminates the use of serial measurements of peak expiratory flow for the diagnosis of occupational asthma. His department receives some monies from grants, donations and legal fees to support the research, but he has no personal financial interest.

REFERENCES

Reference List


