Peak flow rate records in surveys: reproducibility of observers' reports

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ABSTRACT Records of peak expiratory flow rate (PEFR), commonly used in hospital in the management of asthma, have not been evaluated as a method of identifying cases of asthma in population surveys. Four observers were asked to report on whether asthma was present or absent in 61 graphs of PEFR recorded two hourly for four weeks during surveys of working populations. Agreement within individual observers was measured using a subset of 29 graphs which had been copied and distributed at random among the set of 61; agreement was good, from 90% in one observer to 100% in two. Agreement between observers was measured on the basis of all 61 graphs. Agreement occurred between all four observers in 69% of graphs, between at least three out of four in 97%, and, when pairs of observers were examined, between 72% and 93% of graphs. Graphs assessed as showing asthma demonstrated more within day PEFR variability (expressed as the number of days in which the difference between maximum and minimum readings was at least 15%) than graphs assessed as not showing asthma. Some graphs with little within day variability were assessed as showing asthma, apparently because they demonstrated between day PEFR variability.

Asthma, defined as variable airflow limitation,1 can be demonstrated by regular monitoring of lung function, conveniently performed by recording the peak expiratory flow rate (PEFR) over several days or weeks. Such records may be assessed by inspecting the raw readings or graphs drawn from the readings, and this is now common in hospital in the evaluation of the severity of asthma and its response to treatment.2 PEFR recording by patients outside hospital has been encouraged by the introduction of the miniature meter,3 which is extending the use of PEFR records to surveys of asthma in populations.

The widespread acceptance of PEFR records in clinical practice is an endorsement of the method's usefulness. Its validity in diagnosis is, however, difficult to estimate formally, there being no agreed standard test for asthma against which it could be compared. Techniques for identifying disease in epidemiological surveys should be reproducible as well as valid, and for PEFR records one important potential source of variation is differences among observers providing reports on the records. In contrast to hospital practice, records from surveys are assessed in isolation by an observer who is "blind" to other relevant information. Any abnormalities are likely to be minor and difficult to interpret. Variation in reporting might be a serious problem in surveys, as it is when physicians take a history of respiratory symptoms,45 examine the chest,56 or look at chest radiographs.78

We have therefore taken records made during surveys of working populations and measured observer variation in the detection of asthma from these records alone. We have also attempted to identify factors which influenced these observers in their reporting.

Methods

SUBJECTS

Recordings from 61 men formed the basis of the study. Thirty eight men were currently employed in a steel coating plant where isocyanate induced occupational asthma had occurred from 1972 to 1979.
and 23 were employed in an electronics factory where acid anhydride induced occupational asthma had occurred in 1979 and 1980. These PEFR records formed part of follow up studies at the two workplaces in 1982 after occupational exposures had been controlled by, respectively, substitution of another chemical and improved ventilation. On clinical grounds 15 men were originally classified as having asthma related to occupation, 23 as having asthma unrelated to occupation, 20 as having respiratory symptoms not caused by asthma (but by, for example, chronic bronchitis or non-specific irritation), and three as having no respiratory symptoms (but with serum antibody against an acid anhydride-protein conjugate). Many of the subjects, including 12 of the 15 men with occupational asthma, had lost their symptoms or had partially improved by the time the PEFR records were made. We therefore confined this study to the identification of asthma rather than occupational asthma.

**PEFR RECORDS**

Each person was given a mini Wright peak flow meter and instructed in its use. He was asked to note, on a standard form, the best of three readings taken every two hours during waking hours for four weeks. The readings were plotted as previously described as graphs of the maximum, minimum, and mean of each day's PEFR, days at work being indicated by shading. The graph is plotted by computer, which also prints each day's within day PEFR variability, defined as the difference between the day's maximum and minimum readings expressed as a percentage of the maximum. For each record the number of "variable" days, with a difference of at least 15% between maximum and minimum readings, provided an index of PEFR variability.

**REPORTING**

Four observers (the authors) who were experienced in using these records assessed each graph independently and without knowledge of the subjects' identities, symptoms, or exposure. We used a four point scale: 4—definite asthma; 3—probable asthma; 2—probably not asthma; 1—definitely not asthma. This scale was chosen because it offered comments which resembled those made spontaneously on graphs from other surveys and it forced the observer to make a decision on whether asthma was or was not present while recognising the potential difficulties in reporting on graphs from a working population using limited information.

### Table 1 Use of the asthma assessment scale by four observers, reporting on 61 graphs

<table>
<thead>
<tr>
<th>Observer</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asthma</td>
<td>18</td>
<td>20</td>
<td>23</td>
<td>11</td>
</tr>
<tr>
<td>Probable</td>
<td>2</td>
<td>4</td>
<td>8</td>
<td>11</td>
</tr>
<tr>
<td>Not asthma</td>
<td>9</td>
<td>14</td>
<td>3</td>
<td>8</td>
</tr>
<tr>
<td>Definite</td>
<td>32</td>
<td>23</td>
<td>27</td>
<td>31</td>
</tr>
<tr>
<td>Total</td>
<td>61</td>
<td>61</td>
<td>61</td>
<td>61</td>
</tr>
</tbody>
</table>

### COMPARISON OF REPORTS

Twenty nine graphs were taken at random from the set. Copies were made and the copies then returned at random together with the originals. Intraobserver variation was measured by comparing each observer's reports on the 29 duplicates. All 61 graphs were used in measuring interobserver variation and, in the case of the duplicated graphs, the first of each pair was used. Each observer therefore assessed a total of 90 graphs. Complete agreement was defined as the use of the same point on the four point scale. Substantial agreement was defined as either agreement that asthma was definitely or probably present or agreement that it was definitely or probably not present. When used without qualification, "agreement" means either substantial or complete agreement.

### Results

Table 1 shows the different reporting patterns of the four observers and table 2 the intraobserver variation in reporting. These reports agreed in 26–29 (90–100%) of 29 graphs and the agreement was complete in 22–27 (76–93%). Table 3 shows the interobserver variation between all four observers and between the six pairs from the four observers. All four observers agreed on the presence or absence of asthma in 42 (69%) of the 61 graphs and three out of four agreed in a further 17 (28%), so that in 59 (97%) of the graphs most or all the observers were in agreement. For pairs of observers agreement varied from 44 to 57 (72% and 93%) of the graphs.

Figure 1 shows, for the 59 graphs where observers agreed, their assessment that asthma was or was not present compared with the index of within day

### Table 2 Intraobserver variation in the assessment of 29 pairs of graphs

<table>
<thead>
<tr>
<th>Observer</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
</tr>
</thead>
<tbody>
<tr>
<td>Complete agreement</td>
<td>22</td>
<td>24</td>
<td>26</td>
<td>27</td>
</tr>
<tr>
<td>Substantial agreement</td>
<td>4</td>
<td>3</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>Disagreement</td>
<td>3</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>29</td>
<td>29</td>
<td>29</td>
<td>29</td>
</tr>
</tbody>
</table>
Table 3  Interobserver variation in the assessment of 61 graphs

<table>
<thead>
<tr>
<th>Observer combination</th>
<th>All</th>
<th>B/D</th>
<th>B/C</th>
<th>C/D</th>
<th>A/D</th>
<th>A/B</th>
<th>A/C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Complete agreement</td>
<td>29</td>
<td>35</td>
<td>43</td>
<td>39</td>
<td>41</td>
<td>41</td>
<td>37</td>
</tr>
<tr>
<td>Substantial agreement</td>
<td>13</td>
<td>22</td>
<td>9</td>
<td>13</td>
<td>10</td>
<td>10</td>
<td>7</td>
</tr>
<tr>
<td>Majority agreement</td>
<td>17</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Disagreement</td>
<td>2</td>
<td>4</td>
<td>9</td>
<td>9</td>
<td>10</td>
<td>10</td>
<td>17</td>
</tr>
<tr>
<td>Total</td>
<td>61</td>
<td>61</td>
<td>61</td>
<td>61</td>
<td>61</td>
<td>61</td>
<td>61</td>
</tr>
</tbody>
</table>

PEFR variability (the number of days with a difference of at least 15% between maximum and minimum readings). All 22 graphs thought by most or all of the observers to show asthma contained at least one variable day and all nine where observers were in complete agreement that asthma was present contained at least four variable days. None of the 37 graphs thought by most or all of the observers not to show asthma contained more than three variable days and none of the 20 graphs where observers were in complete agreement that asthma was not present contained any variable days. Graphs with one, two, or three variable days were classed both as asthma and as not asthma, and in none of these graphs was there complete agreement between observers.

Figure 2a shows a graph with 16 variable days thought by all observers to show definite asthma and figure 2b a graph with no variable days thought by all definitely not to show asthma. In contrast to these examples, where degree of within day variability was related to the observers’ assessments, are the examples shown in figure 3. Figure 3a shows a graph with two variable days on which there was substantial agreement that asthma was present: it shows a consistent pattern of between day PEFR variability, suggestive of mild work related asthma, rather than within day variability. Figure 3b shows a graph with one variable day on which there was substantial agreement that asthma was not present: it shows no clear and consistent pattern.

In the subgroup of 23 electronics workers an assessment of respiratory symptoms was made at the start of the PEFR record. Of the eight people whose
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graphs were reported by either three or four observers to show asthma, seven experienced current or intermittent wheeze, breathlessness, chest tightness, or unproductive cough and one was symptomless but had serum antibody against an acid anhydride-protein conjugate. Of the 13 thought by either three or four observers not to show asthma, four were symptomless, four complained only of chronic productive cough, four complained of either unproductive cough or intermittent wheeze or breathlessness (none experiencing these symptoms at the time of the record), and one complained of current wheeze and breathlessness but had inhaled corticosteroid and bronchodilator treatment regularly throughout the period of the record.

Discussion

In clinical practice patients referred to hospital with asthma usually have PEFR records showing clear variability within and between days. In contrast, the subjects of this study were working people; the asthmatic subjects had mild asthma and few had consulted a doctor about their symptoms; the others had no symptoms or trivial ones. The fact that physiological changes in this working population were, in general, minor was probably a factor in producing the observer variation we describe.

The reproducibility of the observers' reports was good, particularly the reproducibility of individual observers, who agreed with their own assessments in the great majority of the 29 graphs included in the study of intraobserver variation. Interobserver variation was greater and some may have been semantic, observers differing on the meaning of the assessment scale rather than the meaning of the graphs. This may explain, for example, the pattern of observers B and D, who agreed on the presence or absence of asthma in almost all the 61 graphs but differed on whether it was "definitely" or "probably" present or absent in many. On the other hand, observers A and C disagreed on the presence or absence of asthma in over a quarter of the graphs. Semantic differences may have played a part, or possibly the two used slightly differing criteria for the detection of asthma; or their difference may be explained by differences in experience or personality impossible to evaluate in this study. One of the important functions of studies of observer variation is the stimulation of debate and the development of consensus on criteria for disease detection. We would expect that a second study, similar to this one, with the same observers would show less variation.

As expected, both within day and between day variability of PEFR were criteria used in the detection of asthma. An index of within day variability, the number of variable days, correlated well with the observers' combined assessments, although not necessarily with those of individual observers. In general, records with four or more variable days were regarded as showing asthma and those containing none were not thought to show asthma. Those with one, two, or three variable days could be classed either way, and there was never perfect agreement on the classification of this group of graphs. The use of between day variability as a criterion was most easy to identify in this group, which is exemplified by figure 3a; and this criterion was presumably also used in the interpretation of the other graphs. The pattern of a PEFR graph should be consistent with our knowledge of PEFR variability in health and disease; isolated high readings, such as that in figure 3b, are not and may result from errors in technique or transcription during these unsupervised records. We have noted this previously and
have also noted that PEFR variability in the first
days of a record, while the subject is learning to use
the meter, may be spurious. In clinical practice
doubts about graphs like these can be resolved by
questioning the patient or repeating the record, but
in surveys this is not possible.

PEFR records should be susceptible to a
mathematical analysis and our preliminary work13
(also KM Venables, unpublished cumin analyses)
suggests that the complex and various patterns of
asthma will require a sophisticated analysis for
computer based reporting to be reliable. The human
observer will, however, be necessary, at the least to
test the validity of new mathematical models and to
supplement computer reporting.

We were able to compare symptoms and PEFR
reports in the subgroup of 23 electronics workers
and found a close relationship between the presence
or absence of asthmatic symptoms at the time the
record was made and the PEFR reports made with-
out knowledge of these symptoms.

The epidemiology of asthma is a developing sub-
ject and the PEFR record appears to be a useful
method of identifying asthma. The technique is
non-invasive, inexpensive, and in our experience
acceptable to subjects. We have demonstrated that
observers report reproducibly on the records. We
hope that this report will encourage the use of PEFR
records in surveys in occupational and non-
occupational contexts.

We wish to acknowledge the help given by
Rosemarie Hawkins, who transferred the readings
into a computer file; by David Hughes, who
designed a program for plotting them in a graphical
form; and by Jeanie Thomson, who typed the man-
uscript.

References

1 Scadding JG. Meaning of diagnostic terms in bron-
2 Turner-Warwick M. On observing patterns of airflow
obstruction in chronic asthma. Br J Dis Chest
3 Wright BM. A miniature Wright peak-flow meter. Br
4 Cochrane AL, Chapman PJ, Oldham PD. Observers'
errors in taking medical histories. Lancet 1951;
1:1007-9.
5 Schilling RSF, Hughes JPW, Dingwall-Fordyce I. Dis-
agreement between observers in an epidemiological
6 Fletcher CM. The clinical diagnosis of pulmonary
1952;45:577-86.
7 Fletcher CM, Oldham PD. The problem of consistent
radiological diagnosis in coalminers' pneumoconiosis.
8 Cochrane AL, Garland LH. Observer error in the
9 Dally MB, Burge PS, Davis G, et al. Occupational
asthma in a steel coating plant: population study lead-
10 Howe W, Venables KM, Topping MD, et al. Tetra-
chlorophthalic anhydride asthma: evidence for specific
11 Burge PS, O'Brien IM, Harries MG. Peak flow rate
records in the diagnosis of occupational asthma due to
colophony. Thorax 1979;34:308-16.
12 Venables KM. Two objective methods for the diagnosis
of occupational asthma in epidemiological surveys.
13 Burge PS. Prolonged and frequent recording of peak
expiratory flow in workers with suspected occupa-
tional asthma due to colophony or isocyanate fumes.
Correspondence

8. "This letter was sent to the authors, who reply below.

Sir.—Professor Droszcz and Dr Piotrowska make some extremely important points in their letter. They are correct in the belief that our study was performed on the assumption that there was no difference between the potencies of triamcinolone and triamcinolone acetonide. The investigation was stimulated by the publication of the report in the British Journal of Disease of the Chest in 1979 entitled "Triamcinolone in corticosteroid-resistant asthma." The authors of that study, like ourselves, used doses of Kenalog (triamcinolone acetonide) within the range recommended by the manufacturer in the ABPI Data Sheet Compendium which does not indicate that there is any difference in potency between triamcinolone and triamcinolone acetonide. In experimental animal models it is apparent that triamcinolone acetonide is very much more potent than triamcinolone, but no data from studies in man appear to be available. We therefore have to concede that Professor Droszcz and Dr Piotrowska are perhaps correct in their criticism of the way in which we discussed our data. We think it possible, however, that the information provided by ER Squibb and Sons Ltd about their product Kenalog may have misled the majority, if not all, of the physicians who use this corticosteroid preparation. Although we accept that it is very difficult to assess the relative potencies of corticosteroids, especially when they are administered by different routes, this controversy about the potency of triamcinolone and triamcinolone acetonide highlights the great need for companies to be obliged to state the potency of their products. Perhaps hydrocortisone could be the standard drug with an assumed potency of 1 and the activity of all other corticosteroid preparations for oral, intramuscular, or intravenous compared with it.

If the argument put forward by Professor Droszcz and his colleague about the potency of triamcinolone acetonide is accepted, and so far as we are aware there are no data to refute it, it remains difficult to explain why it causes less suppression of the hypothalamic-pituitary axis (HPA) than daily oral prednisolone in a dose of at least 10 mg. One explanation could be that a large dose of corticosteroid is available very soon after injection of Kenalog, but is not maintained for a full period of four weeks, towards the end of which serum and tissue levels may fall below physiological requirements, with consequent stimulation of the HPA axis. If this is the case treatment with Kenalog could be dangerous when given to patients who have HPA suppression, such as those patients in our study who had been taking large doses of oral prednisolone for a considerable time.

We concluded that we would not normally recommend triamcinolone (meaning triamcinolone acetonide) in preference to prednisolone because of side effects. If triamcinolone acetonide is indeed 10 times more potent than triamcinolone it could never be justified in preference to oral prednisolone in the long term management of bronchial asthma in the doses recommended by the manufacturers. Unfortunately, the data about the relative potencies of triamcinolone acetonide, prednisolone, and hydrocortisone are not published and are only available from ER Squibb and Sons Ltd as confidential information for clinical investigators.

Since publication of our paper we have learned from the manufacturers of Kenalog that it is not an intramuscular depot preparation and the reason for its prolonged but unpredictable duration of action is unknown. We have therefore to admit that even the title of our paper is incorrect.

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Corrections

Peak flow rate records in surveys: reproducibility of observers' reports

In the paper by Dr KM Venables and colleagues (November 1984;39:828-32) we regret that there are errors in the first paragraph of the methods section, in which it is stated that recordings from 61 men were studied. Of the 23 persons employed in the electronics factory, 18 were in fact women. The beginning of the last paragraph of page 828 should read: "Records from 61 subjects formed the basis of the study. Thirty-eight subjects (all male) were currently employed in a steel coating plant... and 23 (18 female) were employed in an electronics factory..." Elsewhere in the paragraph the word men should be taken to indicate subjects.

Bronchial reactivity to inhaled histamine and annual rate of decline in FEV, in male smokers and ex-smokers

Smoking, allergy, and the differential white blood cell count

In the two papers by Dr RG Taylor and others (January 1985) we regret that page numbers are missing from two of the references. In ref 10 on p 16 the pages are 17-22 and in ref 24 on p 21 they are 9-16.