Conventional RIA underestimates cortisol suppression in the presence of prednisolone

Concerns about suppression of the hypothalamic pituitary adrenal (HPA) axis by systemic steroids as well as by inhaled corticosteroids have been widely held since their introduction. Several studies have suggested that inhaled corticosteroids can replace oral corticosteroids during exacerbations of asthma1 and in severe asthma.7 We have recently published a study in which treatment of unstable asthmatic patients for 2 weeks with high doses of inhaled fluticasone resulted in a greater improvement in airway hyperresponsiveness than oral prednisolone.1 At the time of writing this letter we found a comparable decrease in serum cortisol levels with fluticasone 1000 µg twice daily and oral prednisolone 30 mg/day. A radioimmunoassay (RIA) method was used to determine serum cortisol suppression in blood with corticosteroid treatment, as in most studies published to date.6 However, prednisolone and its metabolites are known to be chemically similar to serum cortisol and might therefore interfere with cortisol measurements by RIA.6 Analytical methods involving chromatographic separation of cortisol from prednisolone and its metabolites, such as high performance liquid chromatography (HPLC), circumvent this problem of interference.

We compared serum cortisol measurements by both conventional RIA and by HPLC in the same study,7 which was of a double blind, double dummy, three arm parallel group design. Patients received either oral prednisolone (30 mg/day), fluticasone propionate 1000 µg twice daily (FP2000), or fluticasone dipropionate 1000 micrograms twice daily and oral prednisolone 10 mg once daily in asthma patients. Thorax 1994; 49:37–40.

Figure 1 Change in mean (SE) serum cortisol level (%) from baseline in the three treatment groups measured by conventional RIA and performance liquid chromatography (HPLC).

We conclude that determination of serum cortisol by RIA severely underestimates serum cortisol suppression over a range of 6.9–690 nmol/l serum cortisol in the presence of prednisolone. Our study shows that cortisol suppression in the presence of prednisolone should not be assessed by conventional RIA.

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References

Smoking cessation

We welcome the study by Pelkonen et al1 as a further contribution to our knowledge base on smoking cessation and its effects on pulmonary function and mortality. We feel, however, that some shortcomings in the methodology may bring into question the magnitude of the results.

Our main concern relates to the difficulties in quantifying levels of tobacco exposure. Since tobacco consumption is a continuous variable, confounding factors may occur within each group when categorised too broadly.2 More information about duration and levels of smoking would help to avoid this problem. No information about whether intermittent quitters returned to original habits or resumed smoking at reduced levels. Beneficial effects described in this group could therefore be due to extended periods of decreased tobacco consumption rather than a period of abstinence.

There are no data provided on smoking status from 1974 to 1989. If large numbers of those classed as intermittent quitters had permanently stopped smoking by this time, the value of temporary quitting would be overestimated. Furthermore, no data exist on the duration of periods of abstinence among intermittent quitters. If a significant proportion of this group exhibited prolonged periods of smoking cessation, the relevance of this study to short term quitters is debatable.

Even accepting the beneficial effects of intermittent quitting, we question the importance of this finding in a public health setting. Surely the main health education message must remain the same: permanent smoking cessation should remain the goal and is superior to intermittent quitting. However, we recognise that this finding could provide encouragement to those who have relapsed following
an attempt to quit smoking and reassure them that their failures have not been in vain. This could provide the motivation needed for a second and possibly successful attempt to quit.

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2 Leon DA. Failed or misleading adjustment for confounding. Lancet 1993;342:479-81.

Authors’ reply

Lorna Dunn and Aileen Ogilvie make an important point that the confounding effect of smoking is a strong confounding factor in pulmonary function, which may occur when the levels of tobacco exposure are categorised too broadly. They think that the benefit of intermittent quitting on the decline in FEV\textsubscript{0.75} in our study might be explained by decreased tobacco consumption after periods of abstinence rather than by the periods of abstinence per se. They also point out that, if a considerable proportion of intermittent quitters stopped smoking permanently between 1974 and 1989, it would have led to overestimation of the value of temporary quitting. The third question concerns the duration of periods of abstinence.

In our study the data on smoking habits were recorded at baseline and in subsequent re-examinations by a standard questionnaire. The interval between examinations was usually 3 years. Intermittent quitters were either baseline past smokers who smoked in at least one of the subsequent re-examinations or baseline smokers who were quitters in one or more re-examinations but relapsed back to smoking later. To be recorded as a quitter in an examination a subject had to have given up smoking more than a year previously. During the first 15 years, 27 of 75 intermittent quitters were recorded as quitters in one or more re-examinations (corresponding to at least 1 year of abstinence), 32 were recorded as quitters in two examinations (corresponding to at least 2 years of abstinence), and 16 were recorded as quitters in three examinations (corresponding to at least 3 years of abstinence).

During the first 5 years intermittent quitters reduced the number of cigarettes smoked daily compared with continuous smokers, although not significantly. To measure tobacco consumption more precisely, a new variable was constructed by computing the mean reported daily cigarette consumption at each examination point. For intermittent quitters only, the data from the examinations when they reported smoking were used in making up this variable. When we then additionally adjusted our analyses for this new variable, the decline in FEV\textsubscript{0.75} during the first 15 years was significantly less among intermittent quitters than in continuous smokers (data available from the authors on request). The benefit of intermittent quitting on the decline in pulmonary function therefore also seems to be mediated through periods of abstinence.

Among both intermittent quitters and continuous smokers there were study subjects who stopped smoking permanently between 1974 and 1989. The proportion of such study subjects was greater among intermittent quitters than among continuous smokers. However, when we made additional adjustments for both the mean daily tobacco consumption during the first half of the follow up period and for quitting smoking during the latter half of the follow up period, intermittent quitters still lost less FEV\textsubscript{0.75} during the whole 30 years than continuous smokers (data available from the authors on request).

In conclusion, it seems that some protection may be gained from periods of abstinence, although we agree that the main goal should be permanent smoking cessation.

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- Fibrosing alveolitis in patients with RA

We read with interest the paper by Dawson et al.\textsuperscript{1} on the prevalence of fibrosing alveolitis (FA) diagnosed by HRCT scanning in rheumatoid arthritis (RA). This well-designed cross sectional study estimates the prevalence of FA at 19% in patients with RA irrespective of respiratory symptoms. This is in keeping with current literature and our earlier report of 29% in unselected patients with RA not suspected of having interstitial lung disease (ILD).\textsuperscript{2}

However, neither of these studies has been sufficiently powered to assess a possible association of smoking and FA. Smoking may adversely affect the outcome of ILD in RA and Saag et al.\textsuperscript{3} suspected that smoking was the most consistent independent predictor of ILD patterns in lung function tests and chest radiographs in RA. One of our previous studies\textsuperscript{4} reported a prevalence of ILD of only 5% on HRCT scanning in a cohort of 20 never smokers with RA, while Dawson et al.\textsuperscript{2} reported a prevalence of 11% in never smokers compared with 22% in smokers. There is therefore evidence of a trend towards an association between ILD and smoking which could be explored in a larger study. However, a sample size of 450 patients would be needed to test the hypothesis that smokers are twice as likely to develop ILD in RA than never smokers (95% confidence; power = 80%; smoker/never smoker ratio 2:1).

We agree with the authors that further work on the natural progression of FA diagnosed by HRCT scanning in RA is due. We have commenced a longitudinal prospective study of 18 RA patients with ILD diagnosed by HRCT scanning in a cohort of patients with cryptogenic fibrosing alveolitis (CFA) matched for age, sex, smoking, and respiratory symptoms.\textsuperscript{5} There are significant baseline differences in clinical and radiological features between these two groups. Clubbing and honeycomb appearance on the HRCT scan is more common in patients with CFA while ground glass appearance is more common in RA patients with ILD. The presence of rheumatoid factor appears to be protective against honeycombing in both groups. These differences in clinical and HRCT features may be important predictors of outcome.

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Authors’ reply

We are pleased to receive the letter from Saravanan and Kelly in response to our recent publication in Thorax.\textsuperscript{1} The relationship between smoking and the risk of interstitial lung disease (ILD) in rheumatoid arthritis (RA) is suspected of having interstitial lung disease. We are sure this will provide additional information about the progressive nature of FA and smoking. In the study by Cortet et al.\textsuperscript{6} 68 patients with RA were prospectively studied with HRCT scanning. Cigarette smoking was less prevalent than in the North of England and the ratio of smokers to non-smokers was 1:3. No statistical association was found linking smoking with ILD in RA. A prevalence of 20% of ILD (17% ground glass pattern and 2.9% reticulon pattern) was still found. It is true that in our study the absolute risk of ever smoking cannot be excluded as a risk factor for FA as the number of lifelong non-smokers is small; however, the pack year data are adequately powered to show no statistically significant difference.

With regard to the paper by Rajasekaran et al.,\textsuperscript{7} we feel it necessary to point out that the patients in their study in FA and RA had the diagnosis confirmed by HRCT scanning and, in addition, were symptomatic with dyspnoea, bibasal crepitations, restrictive pulmonary function tests, and chest radiographic changes of FA. We are sure this will provide very valuable information about the progression of FA in patients with RA but it will not add to our knowledge on the outcome of HRCT changes detected at a subclinical stage.

Rajasekaran et al.\textsuperscript{7} found honeycombing on the HRCT scan in three of 18 patients with RA associated ILD and in four of 18 patients with CFA; this difference is not statistically significant.\textsuperscript{8} None of these patients was rheumatoid factor positive, which has led the authors to postulate that rheumatoid factor may be protective against honeycombing. In ILD, these findings may be in direct contrast to those of Muller-Leisse et al.\textsuperscript{9} who found higher levels of rheumatoid factor to be associated with ground glass changes and honeycombing on the HRCT scan, and also to McDonagh et al.\textsuperscript{2} who reported that at least five of 16 patients (31%) had honeycomb and were rheumatoid factor positive. This finding is particularly interesting given that there is evidence in the literature of smoking being associated with seropositivity for rheumatoid factor in patients with RA and without RA.\textsuperscript{10} We...
would suggest that larger studies need to be undertaken and explored for confounding factors such as smoking before a statement can be made that rheumatoid factor is protective against honeycomb.

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References

Measuring granulocyte apoptosis in airway inflammation

We read with interest the paper by Turlej et al describing enhanced survival of lung granulocytes in an animal model of asthma. As discussed by the authors, modulation of immune cell apoptosis is likely to be important in controlling inflammatory processes, and the paper enhances our understanding of this.

However, we feel that there are some methodological problems with the study. Firstly, the animal model they describe, though different from human BLF with no success (unpublished observations) and would be interested to know if the authors achieved this separation easily. We are also surprised at the viability of >90%. Cell viability is likely to diminish with increasing rates of apoptosis, and it is notable that the BAL granulocytes from healthy horses have apoptotic rates of around 40%.

This study is interesting, but the methodological issues raised must be considered in interpreting the results.

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References

Authors’ reply

We thank Dr Kelly and colleagues for their interest in our paper. In the past equine heaves was called COPD but, because equine heaves is very close to atopic asthma and these diseases share important features including hyperresponsiveness to aeroallergens, the Th2 type immune response, chronic airway inflammation, reversible airway obstruction, non-specific airway hyperresponsiveness, and production of specific IgE, it is correct that neutrophils are the predominant inflammatory cells in equine heaves, but this does not exclude the use of this model in asthma studies. Indeed, neutrophils are known to play an important role in these disorders and whereas recent studies have questioned the importance of eosinophils in this disease, we believe that there are only small amounts of granulocytes were recovered from the lung of the horses so we were only able to use one method to assay these cells for apoptosis. We chose the method that has been found to be the most sensitive marker of granulocyte apoptosis—the annexin V (AV)/propidium iodide (PI) method. The results obtained with this method were interpreted as follows: AV−/PI− cells were considered alive, AV+ /PI− cells were considered apoptotic, and AV+/PI+ cells were considered necrotic. This is the first time we have heard of controversy surrounding the interpretation of the results obtained with this method, probably because they have not been published in scientific journals. According to the archives we have read using the web addresses provided by Dr Kelly and colleagues, it appears that this controversy exclusively concerns the status of AV+/PI+ cells. Such cells are uncommon and were not observed in our study.

We agree that density centrifugation may interfere with neutrophil function. To the best of our knowledge there is no other way of separating granulocytes from other cell types. As mentioned in the Methods section of our paper, cell viability of freshly isolated granulocytes was evaluated by trypan blue (TB) exclusion. The cells were then cultured for different times and assayed for apoptosis using AV/PI. Cells in an early state of apoptosis are AV+ and TB−. It is not surprising to find 40% apoptotic (AV+ ) cells in a population where nearly all the cells (>90%) are TB−.

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References

In the Programme and Abstracts of the British Thoracic Society Winter Meeting 2001 published in Thorax 2001; 56(Supplement III), an error occurred in abstract S130 “Management of pneumothorax in a district general hospital” by Al-Aloul M, et al which appeared on page iii40. The name of the second author which should have been KU Toor.

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Measuring granulocyte apoptosis in airway inflammation

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