Bronchiolitis obliterans organising pneumonia associated with the use of nitrofurantoin

R J Cameron, J Kolbe, M L Wilsher, N Lambie

Abstract

The spectrum of nitrofurantoin lung injury continues to widen. The case histories are presented of two patients who developed lung disease associated with the use of nitrofurantoin with histological features of bronchiolitis obliterans organising pneumonia (BOOP), a rare but recognised form of drug induced injury. The two middle aged women presented with respiratory symptoms after prolonged treatment with nitrofurantoin. Both had impaired lung function and abnormal computed tomographic scans, and their condition improved when nitrofurantoin was withdrawn and corticosteroid treatment commenced. The favourable outcome in these two patients contrasts with the fatal outcome of the two other reported cases of nitrofurantoin induced BOOP. We suggest that the previous classification of nitrofurantoin induced lung injury into "acute" and "chronic" injury is an oversimplification in view of the wide variety of pathological entities that have subsequently emerged.

Keywords: bronchiolitis obliterans organising pneumonia; drug induced pulmonary disease; nitrofurantoin

An increasing number of drugs are recognised as causing lung injury and the spectrum of their adverse effects is widening. A recognised but uncommon form of drug induced lung disease is bronchiolitis obliterans organising pneumonia (BOOP).1 We report two cases of nitrofurantoin induced pulmonary disease with histological features of BOOP.

Case 1

A 34 year old female non-smoker with recurrent urinary tract infections presented with increasing dyspnoea and cough over several months. She had been taking nitrofurantoin 50 mg at night for more than two years. She had no other significant exposures and was on no other medications. Examination was normal. The chest radiograph showed diffuse bi-basal reticulonodular shadowing. Baseline blood tests were normal, except for ANA 1:1280, with a wide alveolar-arterial gradient (9.1 kPa).

Carbon monoxide transfer factor (TLCO) was reduced to 67% predicted. High resolution computed tomographic (HRCT) scans of the thorax showed patchy peribronchial interstitial thickening, especially of the medium and small sized bronchi, with very little fibrosis. Open lung biopsy specimens showed that many respiratory bronchioles were distorted and largely occluded by fibroelastic tissue with associated mucus plugging and outgrowth of respiratory epithelium into surrounding alveolar tissue, consistent with BOOP. In the absence of other factors a diagnosis of nitrofurantoin induced pulmonary disease was made and the drug was discontinued. Prednisone 30 mg per day, gradually reducing over nine months, resulted in significant symptomatic improvement, significant improvement in lung function (FEV1, 3.56 l, FVC 4.20 l, TLCO 82% predicted), and considerable but incomplete clearance of interstitial changes on the HRCT scan.

Case 2

A 50 year old female non-smoker with recurrent urinary tract infections gave a two month history of worsening dyspnoea, fatigue, anorexia, and cough with fevers and night sweats for three weeks. There was no history to suggest an underlying connective tissue disorder. She had been taking nitrofurantoin 50 mg at night regularly for one year. On examination she was tachypnoeic and tachycardic with diaphoresis; drug induced pulmonary disease with his-
HRCT scan showed marked reduction of the ground glass opacities and areas of consolidation, but with persistent interstitial fibrosis. Repeat lung function tests showed FEV\textsubscript{1} had improved to 2.88 l (100% predicted) with FVC 2.89 l (77% predicted) and TLCO 66% predicted. The patient was subsequently weaned off oral steroids with no clinical, radiological, or physiological evidence of relapse.

Discussion

We conclude that both patients had nitrofurantoin induced pulmonary disease on the grounds that there was a lack of an alternative explanation for their lung disease and a good response to drug withdrawal and treatment with an oral corticosteroid. We acknowledge that BOOP of other causes may respond well to corticosteroid treatment, but there was no disease recrudescence on steroid reduction and withdrawal. The establishment of a firm aetiological relationship would require re-challenge with nitrofurantoin. This was considered inappropriate in view of the severity of pulmonary impairment on presentation and the residual and irreversible changes on the HRCT scan.

Relatively few pharmaceutical agents have been associated with BOOP. These include amiodarone, acebutalol, nilutamide, cephalosporins, barbiturates, and cocaine. There are only two previously reported cases of BOOP attributable to nitrofurantoin use. Both patients were elderly ex-smokers with symptoms of 3–4 weeks duration and both responded well to initial corticosteroid treatment, but rapid tapering led to an irreversible decline and death after failure to respond to increased steroid dosage. Details of drug treatment were not included in the report.

The course of the disease in our patients was rather different. Both were maintained on medium to high dose prednisone initially, gradually reducing over months, and the duration of treatment may have been important in terms of the improved outcome. They were weaned off oral steroids without clinical, radiological, or physiological evidence of relapse.

Nitrofurantoin induced pulmonary disease may present in many forms including BOOP, diffuse alveolar damage, vasculitis, interstitial fibrosis, pleural and airways disease, and pulmonary haemorrhage. A final common toxic pathway has not been postulated. Nitrofurantoin induced pulmonary disease may result from immune mediated injury or via hydroxyl radical generation with subsequent free oxidant damage. The reduced incidence with the addition of the antioxidant ascorbic acid to nitrofurantoin preparations and results of in vitro studies suggest that this and other antioxidants may significantly reduce toxicity.

Initial reports suggested that the duration of nitrofurantoin treatment dictated the disease pattern. The “acute” reaction was characterised by marked constitutional symptoms including rash, fever, arthralgia, fatigue, together with pulmonary symptoms of dry cough and dyspnoea. The “subacute” and “chronic” forms were more insidious, with increased eosinophil count, raised ESR, and vasculitis and interstitial inflammation on histological examination, consistent with a type III immune response. Increased immunoglobulin levels, hepatic transaminases and ANA titres (the so called “drug induced lupus syndrome”) was associated with a degree of irreversible fibrosis. Some early reports of biopsy specimens from a patient with nitrofurantoin induced lung disease which predate the recognition of idiopathic BOOP as an independent entity are suggestive of a BOOP-like pattern. Cohen suggested that BOOP may be a precursor to chronic lung fibrosis, an early and potentially reversible phase in the spectrum of fibrosing lung disease. However, both patients in this report had residual radiological abnormalities although the remaining functional abnormalities were minor. The subsequent variety of

![Figure 1](http://thorax.bmj.com/)
LETTERS TO THE EDITOR

Systematic review of antistaphylococcal antibiotic therapy in cystic fibrosis

McCaffrey et al  conclude that “antistaphylococcal treatment achieves spu tum clearance of Staphylococcus aureus in patients with cystic fibrosis . . .” and that prophylactic treatment in young children is “ . . . likely to be of clinical benefit” . These positive conclusions are based on the results of a study which has important methodological problems. Neither the introduction nor the methods section of this review state what hypotheses the review set out to test, the criteria used to decide whether a study was suitable for inclusion, outcomes to be studied in the review, or methods used to assess the methodological quality of included studies. Systematic reviews differ from narrative reviews in that they test hypotheses using a methodology which is well described . The authors have described their search strategy, which is based on that developed by the Cochrane Collaboration, to identify randomised controlled trials. The authors have, however, included a number of studies in their review which are not randomised controlled trials. It is not clear from the information provided whether their search strategy is sensitive enough to identify all possible relevant studies.

The authors base their conclusions on the results of just two randomised controlled trials, involving only 66 children, with a maximum follow up of two years . All of these children were under seven years of age (most under two years) and had upper respiratory samples taken, not sputum. Of the other studies described as randomised, one used alternate allocation [and so was not randomised] and one reported further outcomes in patients included in one of the randomised controlled trials. Only two randomised controlled trials actually reported the prevalence of S aureus in respiratory secretions. The larger study by Weaver et al reported that the prevalence of S aureus was reduced with prophylaxis but “clearance” was not achieved from nose and throat swabs. The important issues for cystic fibrosis patients and their families are not eradication of an organism but fewer symptoms, improved lung function, and prolonged survival. None of the studies described in the review addressed these . “This objective is consistent with the view of the authors of the Cochrane Collaboration handbook which recognise that systematic reviews can have different motivations, one of which is the resolution of conflicting evidence.” Indeed, it is probably difficult to define systematic reviews as formally as Smyth et al (and others) have proposed as the science of systematic reviewing is undergoing continuous development. More systematic reviews are being performed now than ever before (a Medline search looking for “systematic review” in titles and abstracts presents 4158 citations in the last 10 years, 1538 (37%) of which are in the last two), with reviewers defining their methods according to the problem in question.

Again, because of the nature of the field being studied, we had purposely not defined stricter criteria for study selection or drawn up a preselected list of outcomes of interest. As the area under investigation was largely unknown, we felt such criteria could limit our search. Also, in the absence of any significant background information, we were uncertain if such a choice of outcomes could be made objectively. Indeed, if we had arbitrarily drawn up a list of outcomes that were of interest to us, we would have missed a number of outcomes that others had used and which could be of potential interest to readers when designing clinical trials in the future. We did not use quality scores because there is little objective evidence to support the use of quality scoring in systematic reviews.

Many of the scoring systems have not been developed with sufficient rigor and could add the analyst’s bias to the results . A recent review of a random sample of 240 meta-analyses showed that less than half assessed trial quality . However, we note that newer techniques such as meta-regression may provide better alternatives in the future.

As we were principally interested in randomised controlled trials (RCTs), we used a search strategy that has been well validated for the recall of such trials. However, as before, we wanted to present an analysis of outcomes of both RCTs and non-RCTs because we felt this would make our conclusions more objective and be more supported by the authors of the Cochrane Collaboration Handbook. Smyth et al state quite rightly that the important issues for cystic fibrosis patients and their families are not eradication of an organism but fewer pathologic entities now shown to be caused by nitrofurantoin suggests that these early categorisations are an oversimplification.

The initial interest in nitrofurantoin induced lung disease has waned as more suitable less toxic agents have been found for chronic urinary infections. However, the drug remains generally available in spite of its high toxic profile and clinicians need to be aware of the spectrum of associated lung disease.


ALAN SMYTH
Children’s Cystic Fibrosis Unit, Nottingham City Hospital, Nottingham NG5 1PB, UK

SARAH WALTERS
Department of Public Health and Epidemiology, University of Birmingham, Birmingham B15 2TT, UK

ROSALIND SMYTH
University of Liverpool Department of Child Health, Royal Liverpool Children’s Hospital, Liverpool L12 2AP, UK


AUTHORS’ REPLY Smyth et al have listed a number of features that they regard as being essential to systematic reviews. In particular, they suggest that systematic reviews should always test hypotheses. However, a clear understanding of the existing evidence is necessary for the generation of valid hypotheses and, in our view, this is best achieved by systematic reviewing. Indeed, many important systematic reviews published in major clinical journals do not specifically test hypotheses, but study the current evidence in order to identify the state of existing knowledge and to define areas for further research. This objective is consistent with the view of the authors of the Cochrane Collaboration handbook who recognise that systematic reviews can have different motivations, one of which is the resolution of conflicting evidence. Indeed, it is probably difficult to define systematic reviews as formally as Smyth et al (and others) have proposed as the science of systematic reviewing is undergoing continuous development. More systematic reviews are being performed now than ever before (a Medline search looking for “systematic review” in titles and abstracts presents 4158 citations in the last 10 years, 1538 (37%) of which are in the last two), with reviewers defining their methods according to the problem in question.

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symptoms, improved lung function, and prolonged survival. However, this should not inhibit the use of laboratory based outcomes which could influence clinical decision making until appropriate clinical data are available. Indeed, given the high predictive value of oral corticosteroids in childhood for identifying pathogens in bronchoalveolar lavage fluid (sensitivity and specificity of 90%),5 we feel the evidence we have defined in support of Staphylococcus aureus from the upper or lower respiratory tract with anti-staphylococcal antibiotics does suggest that this therapeutic intervention is likely to be of clinical benefit, although we strongly support further research to properly designed studies are needed to confirm this hypothesis.

KEVIN MCCAFFERY RICHARD E OLLER MARGARET FRANKLIN SOMNATH MUKHOPADHYAY Centre for Research into Human Development, Ninewells Hospital & Medical School, Dundee DD1 5SY, UK

6 Moher D, Cook D, Jadad A, et al. Assessing the quality of reports of randomised trials: implica-
8 Apostal A, Uswedy K, Picard E, et al. Sensitivity and specificity of oropharyngeal suction versus bronchoalveolar lavage in identifying respira-


Therapeutic ratio of inhaled fluticasone

I read with interest the recent article by Meijer et al on the effects of inhaled fluticasone and prednisolone on clinical and inflammatory parameters in patients with asthma.6 Rather than focusing on the differences between oral and inhaled corticosteroid, I believe that a more important finding is the effect of a fourfold increase in the dose of fluticasone on the therapeutic ratio. For all-way parameters there were no significant differences in the effects on bronchial hyperresponsiveness to adenosine monophosphate or on sputum eosinophils between the two doses of fluticasone in doses of 500 µg and 2000 µg per day. However, for systemic bio-

activity markers there were no significant differences between the two doses of fluticasone on serum corticoster levels and blood eosinophils. Taken together these findings suggest that, at least for effects on airway hyperresponsiveness and inflammation, the therapeutic ratio for fluticasone declines sharply above a watershed dose of 500 µg per day. This result is perhaps not surprising, given the high glucocorticoid topical potency for in vitro anti-inflammatory activity with fluticasone.

It is also important to point out that the study by Meijer et al was performed using fluticasone delivered via a Diskhaler dry powder inhaler device, which delivers a twofold lower respirable fine particle dose than a fluticasone propionate powder in the same dose inhaler.7 This is due to the larger particle size from the fluticasone dry powder inhaler. Hence, increasing the nominal dose of fluticasone dry powder may result in a proportionately greater delivery of larger particles to the central airways and consequently to a less than expected impact on small airway inflammation. The lower fine particle dose of fluticasone dry powder will also result in reduced lung bioavailability, as shown by a fivefold lower degree of adrenal suppression compared with the same nominal dose of fluticasone delivered via a pressurised metered dose inhaler with spacer device.8 The use of fluticasone in a dose of 500 µg/day via a dry powder inhaler would therefore explain the absence of any significant suppression of blood eosinophils or serum cortisol in their study. This does not mean that fluticasone propionate dry powder in a dose of 500 µg/day is not systemic, as Meijer et al recently published data with this dose of fluti-

casone given via a Diskhaler reported significa-

tant suppression of 24 hour urinary cortisol excretion (33% reduction) and peripheral blood lymphocyte glucocorticoid receptor mRNA expression (71% reduction) during steady state dosing in asthmatic patients.

Another finding in the study by Meijer et al was the relatively greater effect on bronchial hyperresponsiveness to adenosine monophosphate than to methacholine challenge with both oral and inhaled corticosteroid after two weeks. Similar findings have been reported after two weeks of treatment with inhaled budesonide powder in a dose of 1600 µg/day.9 The authors not unreasonably suggested that adenosine monophosphate responsiveness might be more sensitive to changes in airway inflammation than metha-

choline. However, the treatment period was relatively short and this cannot exclude the possibility that the effects on methacholine hyperresponsiveness might have been proportionately greater with a longer duration of treatment, as has been reported in previous studies.10 It is also plausible that differences in bronchial hyperresponsiveness between the doses of inhaled fluticasone may have become apparent with a longer duration of treatment.

Finally, it is important not to extrapolate the results of the study by Meijer et al on patients with relatively mild asthma to more severe asthmatic patients in whom altered airway geometry may cause a reduction in lung delivery and lung bioavailability from narrowed peripheral small airways. Also, their results may be specific to the unique drug/device interaction of fluticasone propionate given via the dry powder inhaler, and further studies are needed to look at the dose-response relationship of this therapeutic ratio using more efficient delivery systems such as a pressurised metered dose inhaler with spacer.

BRIAN J LIPWORTH Asthma and Allergy Research Group, Department of Clinical Pharmacology, Ninewells Hospital and Medical School, University of Dundee, Dundee DD1 5SY, UK

1 Meijer RJ, Kerstjens HAM, Arends LR, et al. Effects of inhaled fluticasone or oral predni-
solone on clinical and inflammatory para-

3 Oslo aerosol particle generation from dry-

4 Wilson AM, Dempsey OJ, Coutie WJ, et al. Impotence of drug-device interaction in deter-
mixing systemic effects of inhaled cortico-

6 O’Connor BJ, Ridge SM, Barnes PJ, et al. Greater effect of inhaled budesonide on adenosine-3-monophosphate induced than on sodium metabisulphate induced bronchocon-

8 Haahrela T, Jarvinen M, Kava T, et al. Compari-


Authors’ reply We thank Dr Lipworth for his interest in our article. Although we found no significant dose difference in PC20 adenosine monophosphate and metha-

choline over a two week period between the two doses of fluticasone, the trends suggested a favourable effect of 2000 µg compared with 500 µg per day for every parameter measured, and there was, indeed, a significant dose response effect on sputum levels of ECP. It is well known that the dose response curve for inhaled steroids in general is very shallow at conventional and higher doses, and we agree that from our data this seems to apply to fluticasone also. From our study, in which only two doses of flutica-

sone were used, we are careful not to overinterpret where the decline in the therapeu-

tic ratio starts with this drug.

We are aware that the respirable fraction of fluticasone in the dry powder formulation is lower than in the pressurised metered dose inhaler, although the suggested magnitude of the difference is debatable using data from Dr Lipworth’s own group. Unfortunately, in humans we still have considerable problems in separating the effects of common drugs on the large and the small airways, and the remarks by Dr Lipworth on the site of delivery are intuitively correct but, we believe, unproven as far as the clinical effects are con-
cerned. There is no doubt that the pressurised meter formulation has systemic bioavailability and we clearly demonstrate this. We accept the notion that, with more sensitive markers of bioavailability, an effect might have been demonstrable above the dose used in our study. The clinical relevance of this still needs to be determined even after so many years of using inhaled steroids.

We agree that the improvement in hyperre-

sponsiveness with steroid treatment can con-

tinue for much longer than the improvement in forced expiratory volume in one second (FEV1).11 The concept that the improvement in methacholine hyperresponsiveness might continue for a longer period than that of
adrenosine is interesting, but we are unaware of any data to substantiate this. In fact, in a study by Weersink and colleagues, the same difference between the two bronchoconstrictor agents held true for six weeks instead of the two weeks of fluticasone treatment in the current study. It is interesting to debate whether the insufficient effect of inhaled steroids in patients with severe asthma is due to lower availability of peripheral airways, as Dr. Lipworth suggests, or, for instance, to a decrease in sensitivity to steroids—either per se or as a result of increased inflammation and associated cytokine load. The suggestion by Dr. Lipworth would result in a different shorter effect of systemic steroids compared with inhaled steroids, especially in the more obstructed patients, but this does not agree with our clinical impression. In fact, the finding of a superior effect of the inhaled corticosteroid over oral prednisolone (30 mg for two weeks) in our study rather suggests a contrary mechanism, perhaps compatible with a higher effectiveness of the lipophilic compound fluticasone propionate at the level of the epithelium and (sub)mucotha of systemic prednisolone, even if only in the larger airways. Nevertheless, we are careful not to extrapolate our findings beyond the devices and populations studied. There are, however, in addition to ours, a few other studies which suggest that inhaled corticosteroids may have an effect at least as great as prednisolone in asthma exacerbations.

RONALD J MEIJER HUIJ B A KERSTIJNS DIRKJE S POSTMA
Department of Pulmonary Diseases, University Hospital Groningen, Groningen, The Netherlands


“Systematic review” of asthma education studies

We were disappointed that Sudre et al felt there was insufficient documentation and excessive variability in studies of education programmes for adults with asthma published between 1979 and 1998. We feel that their conclusion is largely because they did not perform a rigorous systematic review of papers in this area.

Systematic reviews of research evidence are undoubtedly invaluable scientific activities. They establish whether scientific findings are consistent and can be generalised across populations, settings, and other variations. Systematic reviews should be based on the “gold standard” of published randomised clinical trials. However, in the 77 trials reported in the review of Sudre et al included 35 studies which were not randomised controlled trials. They also give no information about which interventions were found to have statistically significant effects. They include a study which simply asked patients whether they preferred audio-visual information or written information and did not have any intervention, a study which has not been published, and interventions assessing the use of psychotherapy and yoga for asthma patients, which seem well outside the criteria for inclusion in the review. Another four studies they include are from the Cochrane reviews of patient education on the grounds that they are not educational interventions at all. It is not surprising that in 81% of projects assessed the background educational theory was not mentioned and few projects had a patient’s needs assessment performed.

While we accept that many of the studies reviewed had missing information on the form and duration of education, we are concerned that some of these studies may be being misquoted. As an example, our own randomised controlled trial on personalised patient education for asthma delivered in four booklets over three months (reference 65) is described in sufficient detail what that intervention consisted of. We therefore included in our review all studies that had an educational component, regardless of the evaluation design.

We admit that we used a broad definition of education as “any attempt to provide the patient with knowledge or personal skills to reduce the impact of asthma.” The educational content varied among programmes (this is one of our main points) and could include drug management, environmental control, relaxation, yoga, etc. The paper by Partridge et al described an educational intervention without specifying the description of an education programme in an asthma clinic, its weaknesses, and attempts at correcting these. As for including work published only as a dissertation, this may be considered an advantage rather than a drawback by some meta-analysts. We maintain that all studies that we reviewed included an explicit educational component and doubt that changing eligibility criteria to exclude a small subset of studies would much alter our general conclusions.

We stood corrected about the incomplete reference to the Grassic intervention in the discussion section of our paper. In our original base programme this was described more accurately as follows (partial data): number of training sessions: 4 (counting one 10 minute session in person and three mailed booklets); duration of training period: 3 months; delivery of education by: physician and self-help; educational setting: individual; training tools: booklet; training method: lecture/vertical teaching. Had we conducted an effectiveness review we would have noted that the Grassic single out this study as by far the largest trial of asthma education, and one that did achieve clinical benefits for its patients. More such research studies are needed.

BOOK REVIEWS


This text is a thorough but concise overview of clinical tuberculosis presented as a well structured series of cases with clearly reproduced radiographs, computed tomographic scans, and slides. Each case is complemented by a short pertinent discussion clarifying any points of interest or debate. A carefully chosen chapter layout sequentially introduces the reader to the most challenging and interesting aspects of the disease and also provides an easy reference framework.

The authors’ obvious wealth of experience allows readers with a more limited exposure to learn something of the more unusual manifestations of infection, including an extensive range of extrapulmonary and multisystem disease. The complex matter of antituberculous treatment in the emergent group with drug resistant mycobacterial infection, comorbidity, or compliance problems is tackled in some depth, highlighting potential pitfalls and explaining, in a real clinical context, the reasons behind the decisions made.

The difficulties associated with the diagnosis and management of tuberculosis in patients with human immunodeficiency virus are well illustrated, but not exhaustively covered, in a chapter whose commentary sections are particularly full and instructive.

Most of the 120 featured case presentations have a short list of affiliated references aimed to guide, rather than delineate in detail, further research of the points of interest raised.

The format of the book ensures an enjoyable and pragmatic approach to learning about tuberculosis, thus making it directly relevant to all those involved in the medical care of patients with the condition, especially at a training level. It would be an ideal accompaniment to existing formal textbooks.—ILJ


This is one of a series of publications under the collective heading “Progress in Inflammation Research” to which some of the European heavyweights in asthma research have contributed chapters. All the asthma drugs are included with the notable exception of the anticholinergic agents, although I found the title a little misleading as the in vivo anti-inflammatory effects of some of the drugs discussed remains contentious. However, from the opening chapter it becomes apparent that investigations into the pathophysiology of, and the effects of treatment on, asthma have played an important part in defining the inflammatory mechanisms. The “commonly” used asthma medications are discussed initially with Peter Barnes giving an erudite synopsis of the anti-inflammatory effects of corticosteroids. The next two chapters deal with the putative anti-inflammatory effects of phosphodiesterase inhibitors and β2 adrenoceptor agonists, although the chapter on phosphodiesterase inhibitors concentrated on the different isoenzymes and thus was heavy going with little discussion of their anti-inflammatory effects and no concluding summary. Despite theophylline being available for at least 40 years, I was struck by the paucity of clinical data available regarding its efficacy and in vivo anti-inflammatory effect (if at all). This is presumably because it is not profitable for pharmaceutical companies to investigate the drug further. The mast cell stabilisers are considered next, and the last third of the book deals with leukotriene antagonists and discusses other novel potential anti-inflammatory agents including anti-IgE agents, cytokines and adhesion molecule antagonists.

Several of the chapters are interesting and well written with well laid out tables and graphs, although some have several annoying typographical errors. The book does provide a good summary of the anti-inflammatory effects of present and potential future asthma medications and would act as a good reference source for departments or individuals with an interest in this field.—JB

NOTICES

Cardiovascular Disease Prevention V

A conference entitled “Cardiovascular Disease Prevention V” will be held on 4–7 April 2000 at the Conference Centre, Kensington Town Hall, London. For further information contact Haringey Medical Conferences Ltd, 127 High Street, Teddington, Middlesex TW11 8HH, UK. Telephone +44 (0)181 977 0011. Fax +44 (0)181 977 0055. email hmc@hamptonmedical.com

British Association for Lung Research

The British Association for Lung Research (BALR) Spring Meeting entitled “Inflammation Control: A Goal for the Millennium” will be held on 18 April 2000 at the Wills Hall, University of Bristol. For further information contact Dr Lynne Armstrong, The Lung Research Group, University of Bristol Medical School Unit, Southmead Hospital, Westbury on Trym, Bristol BS10 5NB, UK. Telephone +44 (0)117 959 5348. Fax +44 (0)117 959 5018. email Lynne.Armstrong@bristol.ac.uk

UK Pulmonary Vascular Units

In the list of UK Pulmonary Vascular Units given at the end of the review article on “Primary pulmonary hypertension” by A J Peacock which appeared in the December issue of Thorax (1999;54:1107–18), the address for Dr Simon Gibbs should have included the Imperial College School of Medicine which includes Hammersmith, Brompton and Harefield hospitals.

Atrial septostomy in pulmonary vascular disease

In the editorial entitled “Role of atrial septostomy in the treatment of pulmonary vascular disease” by R J Barst which appeared on pp 95–6 of the February issue of Thorax, there was an error in figure 1. The correct version is reproduced below, showing that in “non-responders” the PAP is increased or unchanged. The publishers apologise for this error.

CORRECTIONS

Recurrent syncope and/or right heart failure with intact atrial septum

Cardiac catheterisation

Acute vasodilator drug testing

"Responder" (PAP or PAP unchanged)

"Non-responder" (PAP or right heart failure with intact atrial septum)

Continuous PGI2

Improves

Chronic oral vasodilator treatment

Improves

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