Proceedings of the British Thoracic Society

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Spatial mapping to relate respiratory morbidity to a point source of air pollution

P MILLIGAN, VJ KELLY, BJ BRADBURY, REID, E DUNNE, MG PEARSON School of Tropical Medicine, Sefton Health, Sefton MBC and Aintree Chest Centre, Liverpool. Previously we reported a 60% increase in excess cough and significantly increased absences from school due to respiratory symptoms, in an area of increased dust pollution in a survey of 1872 children aged 5–11 years in Merseyside. We extended the analysis to relate levels of airborne dust pollution, respiratory symptoms and distance of residence from a point source of air pollution.

Residential locations were mapped using postcode data. Airborne dust pollution was monitored for 12 months using dust deposition gauges in the Liverpool dock area and two other areas of the city thought to be typical of urban pollution. Mean dust deposition levels decrease radially as distance from the dock area increases; 0.5 km, 115 mg/m²/day; 1.0 km, 90 mg/m²/day; 2.0 km, 40 mg/m²/day.

Children living within a 1 km radius of the dock area have a twofold increased risk of respiratory symptoms than those 3 km from the dock (OR 2.1, 95% CI 1.5 to 3.0) after adjusting for other environmental, predisposition and socioeconomic risk factors. For children within 1 km 35% had excess cough and 24% breathlessness against 18% and 11% respectively more than 2 km away. The control areas had 16% and 19% excess cough respectively. There is a relationship (p<0.05) between mean dust pollution levels in the vicinity of the child’s residence and the risk of having excess cough which halved at a distance 2 km from the dock. The technique of spatial mapping can help identify a probable point source of pollution even in a cross sectional study.

Aeromycology of hospital buildings: non-construction related aspergillosis and detection of contaminated aerosols

G MORRIS, H KENNEDY, J BUTCHER, G SHANKLAND, M RICHARDSON, K ANDERSON Departments of Respiratory Medicine, Medical Microbiology, and Bacteriology, Western Infirmary, Glasgow and Public Health, University of Glasgow. Nosocomial aspergillosis has been generally associated with building construction in close proximity to immunosuppressed patients. While this observation is likely to be correct for some infection episodes, large hospital sites usually have a continuous programme of building alteration, and there is recent evidence to suggest that building demolition or construction is not always followed by measurable differences in aeromycology (Goodeley et al. J Hosp Infect 1994). We recently investigated two separate nosocomial outbreaks of aspergillosis which developed in a paediatric oncology ward (five cases over six months) and adult haematology ward (seven cases over nine months), neither associated with a discrete spell of construction or demolition. Although both units were served by filtered air conditioning, non-filtered air could enter through windows or the hospital waste disposal conduit to the basement of the multi-storey building design. A fumigatus was isolated from pigeon excreta close to the windows in the adult unit, and was also consistently isolated in air and swab samples around the waste conduits in both units (as well as A niger, flavus, terreus, versicolor and other species – predominately Penicillium). A fumigatus was isolated within the paediatric unit vacuum cleaner and its exhaust (65 colony forming units/m³ to 0–6 cfu/m³ background) which possibly acted as a biological trap and disseminator. The adult unit vacuum cleaner was also contaminated but this reflected the ward predominant airborne organism of Penicillium species. Both outbreaks have resolved after sealing of the disposal conduits and removal of the contaminated vacuum cleaner. Neither of these routes have been reported as a source of nosocomial aspergillosis previously and underscore the importance of all sources of air and aerosols close to this group of patients.

Employment status and the perceived association of occupation with respiratory symptoms in young adults with and without asthma

D JARVIS, P BURNEY, S CHINN, C LUCZYNSKA, E LAI, R HALL, B HARRISON, J STARK Department of Public Health Medicine, UMDS; Department of Respiratory Medicine, The Ipswich Hospital; Department of Respiratory Medicine, West Norfolk Hospital; Department of Respiratory Medicine, Addenbrookes Hospital, Cambridge. As part of the British arm of The European Community Respiratory Health Survey data on respiratory symptoms and occupational history were collected from a random sample of young adults living in East Anglia. Subjects were considered to have asthma (ast) if they reported being woken by shortness of breath in the last 12 months or asthma attacks in the last 12 months

<table>
<thead>
<tr>
<th>Current employment status (%)</th>
<th>Student</th>
<th>Employed</th>
<th>Unemployed</th>
<th>Housewife</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asthma (n=184)</td>
<td>3.6</td>
<td>73.8</td>
<td>11.8</td>
<td>10.8</td>
</tr>
<tr>
<td>No asthma (n=1007)</td>
<td>3.3</td>
<td>80.4</td>
<td>6.5</td>
<td>9.8</td>
</tr>
<tr>
<td>Being at work makes chest tight or wheezy</td>
<td>Asthma</td>
<td>33.2%</td>
<td>8.6%</td>
<td>45.4%</td>
</tr>
<tr>
<td>Ever exposed to vapours, dust, gas, fumes at work</td>
<td>55.3%</td>
<td>45.4%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Changed job because affected breathing</td>
<td>6.5%</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Respiratory symptoms and asthma in a working population and in the general population

SC STENTON, JR BEACH, G DEVEREUX, SJ BOURRE, DJ HENDRICK Chest Unit and Regional Unit for Occupational Lung Disease, Newcastle General Hospital, University of Newcastle upon Tyne. We recently performed two investigations of asthma prevalence involving 876 subjects selected at random from FHSA registers (20–44 years, 289 women) and 1126 shipyard workers (16–25 years, 50 women). All subjects completed a respiratory questionnaire and underwent measurements of airway responsiveness as PD20-FEV1, to methacholine. The distribution of PD12 measurements was similar among the two populations as was the proportion in each PD20 category with a physician-made diagnosis of asthma (table 1). The overall prevalence of asthma symptoms was higher among the general population than among the workers. The general population sample also showed a close association between...
symptom prevalence and PD20 category, but the workers showed little difference in symptom prevalence with changing PD20 (table 2). The questionnaire was thus a poorer guide to the presence of asthma among the workers than among the general population. This might have been due to underreporting of symptoms by some workers worried about job security, and overreporting by others concerned about occupational exposures. Similar biases could distort the results of asthma questionnaires under other circumstances.

### Occupational asthma in East London

SK MEREDITH AND P CULLINAN Department of Epidemiology and Medical Statistics, London Hospital Medical College at QMW, London; Department of Occupational and Environmental Medicine, National Heart and Lung Institute, London To assess the aetiological role of occupation in asthma, a postal survey was undertaken of all asthmatics, 16–64 years old, registered with seven general practices in East London (1430 persons). The questionnaire requested the age of onset (or exacerbation) of asthma symptoms and detailed occupational and smoking histories; home visits were conducted on a 20% random sample of non-responders. A net response of 72% was obtained. The prevalence of registered asthma was 4.4%, slightly higher in women, with little difference by age. Half of the men (49%) and 64% of the women reported onset of asthma after the age of 16 years; of those with childhood onset, 41% reported a worsening in adulthood (35% of men, 47% of women). A case-control analysis was carried out of the occupations at the time symptoms began for the 658 persons who reported an onset or exacerbation of asthma after leaving school. Cases (n = 186) were defined as those whose asthma began <2 years after starting a new job and controls as those whose asthma began while in work, but >2 years from starting a job. High risk occupations were defined as those with at least five times greater risk of occupational asthma than the general working population (SWORD data). No association was found in women between starting a high risk job and onset of symptoms. In men the odds ratio (OR) was 1.2 (95%CI 0.5 to 3.3) overall, but was higher in men under the age of 45 years (1-9) and in smokers (1-7). The increased risk was confined to those with childhood onset and exacerbation in adulthood (OR 5-3). The attributable proportion derived from these data suggest that about 2% of asthma in working men in East London is associated with high risk work.

### An in vitro biological effect of an antibody to an epithelial cell autoantigen in cryptogenic fibrosing alveolitis (CFA)

WAH WALLACE, D LAMB, SEM HOWIE Department of Pathology, Edinburgh University Medical School, Teviot Place, Edinburgh EH8 9AG We have previously described the presence of circulating IgG autoantibodies to an extractable lung antigen of 70–90 kDa (Thorax 1994;49:218–24) in patients with CFA. Using serum obtained by immunisation of a rabbit with the partially purified antigen we have demonstrated that it appears to be lung specific, associated with alveolar epithelial lining cells, and present in the A549 type II epithelial cell line (Thorax 1994;49:1139–45). In order to assess whether this antibody might have biological activity we cultured A549 cells and a pulmonary fibroblast cell line in medium containing a range of concentrations (0.001–10%) of immune or control rabbit serum for 72 hours in 96 well plates. The number of cells in each well at the end of this period was assessed using an MTT assay (J Immunol Methods 1983;65:55–63). The results indicated that the immune rabbit serum at concentrations between 3% and 0.1% resulted in significantly fewer A549 cells after 72 hours than the presence of identical concentrations of control rabbit serum (3% p<0.05, 1% p<0.01; 0.5% p<0.001, 0.1% p<0.05). At concentrations lower than this the no differential effect was observed. Further experiments conducted using purified control and immune rabbit IgG in the presence of fresh and heat inactivated rabbit serum confirmed that the effect on the A549 cells was IgG and complement mediated. The rabbit antiserum directed against the putative 70–90 kDa antigen in CFA appears to have in vitro biological activity against the A549 cell line in vitro.

### Epstein-Barr virus (EBV) in type II pneumocytes in cryptogenic fibrosing alveolitis

J EGAN, JP STEWART, PS HASLETON, JR ABBARD, KB CARROLL, A WOODCOCK North West Lung Centre, Department of Histopathology, Wythenshawe Hospital, Manchester; Department of Molecular Biology, Paterson Institute for Cancer Research, Christie Hospital Manchester Cryptogenic fibrosing alveolitis (CFA) is a clinically heterogeneous condition. Both environmental and infective factors have been implicated in its aetiology. We have investigated surgically obtained lung tissue from 20 CFA patients (17 men, mean age 58 years, range 39–69) for evidence of EBV replication and compared this with lung tissue of 21 control patients (n = 21, 14 men, mean age 61 years, range 32–75). Only patients with a typical histological and clinical picture were designated CFA. Fourteen of the 20 CFA patients had received no specific therapy for CFA at the time of biopsy. Monoclonal antibodies directed against the EBV viral antigens, EBV viral capsid antigen (VCA), EBV viral capsid antigen (VCA), MAB 817; Chemicon International Temecula, California, USA) and gp340/220 antigen, which are expressed during the lytic (replicating) phase of the EBV life cycle, were studied. Expression of viral and cellular markers within the tissue blocks were analysed by immunofluorescence using a streptavidin-biotin amplification step. Fourteen of 20 (70%) of the CFA patients were positive for both EBNA (anti-EBNA IgG; >1:50) and EBV antigens (of 21 (9%) of the control group (X2 test, p = 0.0002). In patients with positive staining for EBV viral replication was localised to pulmonary epithelial cells using the specific epithelial cell marker (m613) and immunoperoxidase staining confirmed the staining to be within type II alveolar cells. This is consistent with the first report of EBV replication within the epithelial cells of the lung of AIDS patients in an immunocompetent host. This suggests that EBV may be an immune trigger or contribute to lung injury in CFA, offering a new avenue of treatment.

### Th2 lymphocytes predominate in the pulmonary interstitium of patients with cryptogenic fibrosing alveolitis (CFA)

WAH WALLACE, BA RAMAGE, D LAMB, SEM HOWIE Department of Pathology, Edinburgh University Medical School, Teviot Place, Edinburgh EH8 9AG CFA is characterised by a persistent inflammatory process in the lung which is believed to be immunologically mediated. As such the CF4 helper/inducer lymphocyte subset is likely to play a key role in the disease process. CD4 positive lymphocytes can be divided into Th1 and Th2 subtypes on the basis of the profile of cytokines which they produce (J Immunol 1986;136:2348–57). Cells associated with delayed type hypersensitivity reactions (Th1) are characterised by the production of IL-10 and INFγ, while those associated with the development of allergic type antibody responses are characterised by the production of IL-4 and IL-5. We examined lung biopsy specimens from 10 patients with CFA [median age (range) 70 (61–78)] using a non-isotopic in situ mRNA hybridisation technique (Exp Dermato 1992;1:230–5) and immunohistochemistry in order to identify cells positive for the key cytokines INFγ, IL-4, and IL-5. In situ hybridisation revealed that the majority of infiltrating cells with
lymphocyte morphology contained detectable mRNA for each of the 3 cytokines. In contrast to the immunohistochemical data revealed that while most cells were positive for IL-4 and IL-5 only a very few were positive for INFγ. This difference was quantified using an image analysis system (HOME) [median % lymphocytes positive with antibody (range): IL-4 78% (53-85); IL-5 76% (55-83); INFγ 5-6% (2-13)]. The methylation pattern between detectable mRNA and protein production has been previously noted in other situations (Immunology Today 1990;11:458-64) and suggests that care must be taken in interpreting data on cytokine production when only mRNA is measured. CFA is associated with a predominantly Th2 pattern of CD4 activation.

Pulmonary vascular endothelial permeability in systemic sclerosis: relationship to alveolar epithelial permeability and disease extent

JB CAILES, AU WELLS, R LANEY, CM BLACK, DM HANSELL, R COKER, RM DUBOIS Royal Brompton Hospital, London In patients with systemic sclerosis (SSc) endothelial abnormalities have been found in areas of histologically normal lung, suggesting endothelial abnormalities may precede fibrosing alveolitis (FA). Increased alveolar epithelial permeability is also seen in SSC, and is associated with subsequent lung function decline. We aimed to assess the relationship between endothelial permeability and both extent of FA and epithelial permeability in patients with SSc. Twenty-two non-smoking patients with SSc (17 women) of mean (SD) age 50 (10.8) years were randomly selected. All underwent estimation of endothelial and epithelial permeability during one week. Endothelial permeability was assessed by measuring the pulmonary transvascular flux of 99mTc-labelled transferrin corrected for changes in pulmonary blood volume using 99mTc-labelled red blood cells. Epithelial permeability was measured by the clearance of nebulised 99mTc-DTPA from the lung. The lung was divided into three zones (upper, middle and lower) and endothelial and epithelial permeability was assessed in each. Nineteen of the patients underwent concurrent CT scans (3/10 mm sections). Visual estimates were made by two observers at four anatomical levels of the percentage of abnormal lung (to the nearest 5%) and combined to give an overall percentage of abnormal lung. Pulmonary endothelial permeability was significantly related to epithelial permeability in the lower lung zones only (r=0.58, p=0.005), no relationship being present in the mid and upper zones. Radiological abnormality was not associated with endothelial permeability as there was no relationship between CT extent of abnormality and endothelial permeability in any lung zone, and upper zone endothelial permeability in nine patients with no evidence of upper zone disease on CT scan was no different from patients with evidence of disease. Endothelial permeability did not differ regionally throughout the lung whilst, in contrast, both the extent of CT abnormality and the epithelial permeability were greater in the lower zones than the upper. We conclude that in SSc changes in pulmonary endothelial and epithelial permeability are associated only in the lower zones of the lung where fibrosis is maximal. In the lower zones of the lung the epithelial and endothelial abnormality may predispose to the development of fibrosis. [Supported by a grant from the Raynauds and Scleroderma Association UK.]

Cryptogenic fibrosing alveolitis with normal lung volumes

MJ DOHERTY, MG PEARSON, PMA CALVERLEY, Aintree Chest Centre, Fazakerley Hospital, Liverpool Cryptogenic fibrosing alveolitis (CFA) typically presents with a restrictive pulmonary defect and a low TLCO. However, relative preservation of lung volume has been reported in CFA patients with concomitant emphysema (Turner Warwick. Respir Med 1990). How frequently this occurs and its relationship to symptoms and smoking intensity is not known. We have therefore retrospectively analysed all patients with CFA who attended our laboratory for full pulmonary function tests in the last three years. CFA was the physician diagnosis based on clinical examination, chest radiography, lung function tests and CT scans (28 cases) excluding cases associated with connective tissue disease and asbestos. There were 41 patients of whom 20 had a % predicted TLC of more than 80% (group A) and 21 less than 80% (group B). Group A patients were more likely (χ² test, p=0.05) to be male (75% v 38%), to have finger clubbing (65% v 28%), and to present with non-respiratory symptoms (25% v 0%). They were more likely to be smokers or ex-smokers and had smoked more (40 v 29 pack years). Self reported exercise tolerance, age (61 v 63 years) and time to diagnosis were similar in each group, as was TLCO; however, KCO was significantly lower in group A patients. Group A were more likely to have emphysema reported on CT scan (7 of 12 v 5 of 16 scans) but the difference was not significant. In our laboratory CFA often presents without substantial volume restriction. Whether this reflects a different form of the disorder, a different stage of the disorder or an interaction with emphysema needs further elucidation.

Rapid clearance of inhaled 99mTc-DTPA independently predicts lung functional decline in fibrosing alveolitis associated with systemic sclerosis

JB CAILES, AU WELLS, CM BLACK, R UNDERWOOD, RM DUBOIS Royal Brompton Hospital, London In patients with fibrosing alveolitis (FA) increased clearance of inhaled technetium-labelled diethylene triamine pentaacetate (99mTc-DTPA) is a risk factor for decline in lung function, and the predictive value of the test is increased if two measurements are made. It is not known if the 99mTc-DTPA scan is independently predictive when demographic, disease and lung function variables are taken into account. This study aimed to assess the value of two 99mTc-DTPA scans for prediction of lung function decline, and to determine if the predictive value was independent of other variables. Fifty-four non-smoking patients (39F/15M), with systemic sclerosis and fibrosing alveolitis (FAScS), mean (SD) age 48 (11-7) years had two 99mTc-DTPA scans performed a median of 12 (9-16) months apart. A clearance half time of less than 40 minutes was regarded as abnormal (A) and greater than 40 minutes normal (N). Two groups were defined on the basis of the two scan results:

- Group 1, NN (n=13) or AN (n=7); and group 2, AA (n=34). Survival analysis (life table) was utilised with an event defined as a 15% or greater fall in either forced vital capacity (FVC) or gas transfer (TLCO) from the time of the second scan. Proportional hazards regression analysis was utilised to assess the independence of 99mTc-DTPA clearance in prediction of decline in lung function. Only two of 20 patients in group 1 experienced a significant fall in lung function when compared with 17 of 34 in group 2, p=0.001 (log rank test). The risk of a significant fall in lung function in group 2 remained highly significant (p=0.007) when controlling for the effects of age, sex, past smoking history, initial lung function (TLCO), type of systemic sclerosis (limited v diffuse disease) and treatment. The estimated hazard for lung function decline for patients in group 2 was 10-9 times that in group 1. Abnormal clearance of 99mTc-DTPA on two occasions independently predicts a markedly increased risk of lung function decline in patients with FASc. [Supported by a grant from the Raynauads and Scleroderma Association UK.]

Relationship between HIV-1 proviral DNA copy number in alveolar macrophages and clinical disease in the lung of AIDS patients

JR CLARKE, AJ GATES, RJ COKER, JD WILLIAMSON, DM MITCHELL Department of Medical Microbiology, St Mary's Hospital Medical School, London and the Department of Virology and Immunology, St Mary's Hospital, London Analysis of HIV-1 proviral DNA copy number in cells from the lung of AIDS patients and to investigate possible correlation with clinical disease. A total of 66 AIDS patients undergoing fibrobronchoscopic copy for new respiratory episodes were studied. Alveolar macrophages (AM) were purified by adherence to plastic culture plates. Quantitative polymerase chain reaction (PCR) of HIV proviral DNA sequences was carried out using two different methods; an in house radiometric method and the Amersham Quanti Amp scintillation proximity assay.
Samples from 15 patients were also tested by limited cell dilution. The mean HIV-1 proviral DNA copy number was higher in the AM of patients where a specific opportunistic pathogen had been recovered from the lung ranging from 910 to 1169 HIV copies/10^6 AM compared to 281 HIV copies/10^6 AM in individuals where no respiratory pathogen was isolated. Median HIV copy number in AM was found to be lowest in individuals undergoing their first respiratory episode (10 AM) compared to patients with recurrent respiratory disease (200 copies/10^6 AM). Sequential samples taken from 6 patients with repeated episodes of respiratory disease showed that there was an increased HIV DNA copy number in patients predicted to be at risk of HIV pulmonary disease is lowest when no pathogen can be recovered from the lung although this difference was not statistically significant (p>0.1). HIV proviral DNA copy in AM increases with disease progression.

Quantity of HIV-1 proviral DNA in alveolar cells may be contributing to abnormalities of lung physiology in AIDS patients

JR CLARKE, AJ GATES, BJ COKER, JD WILLIAMSON, JM MITCHELL Department of Medical Microbiology, St Mary's Hospital Medical School, London Measurement of transfer factor for lung carbon monoxide (TLCO) has been shown to be a sensitive method for the detection of respiratory physiological abnormalities in HIV seropositive individuals in the absence of overt lung disease. Sequential testing of lung function in asymptomatic HIV-1 seropositive patients showed that individuals with a TLCO value of <80% predicted normal progressed significantly faster to AIDS than individuals >80% TLCO value. Previously we have reported that mean TLCO values were lower at presentation in patients with more advanced HIV disease compared to asymptomatic individuals. In this study we have used quantitative PCR to investigate the relationship between HIV DNA copy number in BAL and TLCO. HIV-1 proviral DNA was detected by PCR in the bronchoalveolar lavage (BAL) cells of 82 out of 124 (66%) HIV seropositive individuals undergoing diagnostic fiberoptic bronchoscopy for respiratory disease. Successful recovery of infectious HIV-1 from BAL cells by co-cultivation was achieved from 58 out of 100 (58%) individuals indicating that, in the majority, HIV-1 is replication-competent. Detection by PCR or isolation of HIV from BAL cells alone was not related qualitatively to TLCO values in this patient group. However, quantification of HIV proviral DNA by PCR techniques revealed that both mean and median DNA copy number increased from a mean (SD) of 10 (13-5) and 7-5 (median) in individuals with a TLCO value of greater than 70 predicted normal to 625 (1088) and 100 DNA copies per 10^6 BAL cells in patients with a TLCO of between 40 and 49 predicted normal. The highest HIV proviral DNA copy number (1800 (20010) median) DNA copies per 10^6 BAL cells was observed in individuals with a TLCO of less than 29 predicted normal. These differences in the quantity of HIV proviral DNA were statistically significant (p<0.01). We conclude that there is a relationship between HIV-1 load in the lung and abnormal pulmonary physiology of AIDS patients.

Regional microbiology of the lung in patients with cystic fibrosis

DL SMITH, EG SMITH, DE STABLEFORTH, ME KAUFMANN, TL PITT Adult Cystic Fibrosis Unit, Birmingham Heartlands Hospital; Central Public Health Laboratory, London Recent applications of genotyping methods to distinguish strains of Pseudomonas aeruginosa (Pa) and Pseudomonas cepacia (Pc) in the sputum of cystic fibrosis (CF) patients have provided insight into the frequency of multiple strain carriage within individuals that may relate to the interpretation of epidemiological investigations. We have investigated multiple strain carriage by sampling directly from the post-mortem lung. Swabs were obtained from both lungs and the major airways of five CF patients at post-mortem. Pa isolates from these swabs were genotyped using the D2 A gene probe and CHEF typing. Ninety percent of swabs produced successful cultures. Four patients grew Pa from all sites cultured, one patient grew Pa from two lobes and the major airways and Pc from all lobes and the major airways. In four patients, including the patient with co-existent Pc infection, all Pa isolates for each individual were indistinguishable by both genotyping methods. In addition Pa isolates from two of these patients were indistinguishable. In one patient three genotypically distinct strains of Pa were found in different anatomical sites; not all were reflected in major airway cultures. The finding of co-existent Pa and Pc infection in one patient does not accurately reflect the microbiological flora present in the lungs. One of five patients studied was found to exhibit multiple strain carriage which was only partially reflected by major airway culture. The finding of genetically indistinguishable strains in two patients supports the possibility of patient to patient transmission as a source of infection. In general, our study supports the notion that major airway sputum accurately reflects the microbiological flora present at more peripheral sites. However, we have shown that in cases exhibiting multiple strain carriage not all strains present in the periphery may be reflected in major airway cultures.

β2-integrin expression on circulating blood neutrophils from patients with cystic fibrosis

D O’RIORDAN, J HAYES, C O’CONNOR, MX FITZGERALD Department of Medicine, University College Dublin and St Vincent’s Hospital, Dublin Lung disease is the major cause of morbidity and mortality in cystic fibrosis (CF). Lung destruction results from chronic pulmonary inflammation, characterised by the continual influx of neutrophils from the bloodstream. Recently, hyperexpression of the β2-integrin (CD18) adhesion molecules on circulating neutrophils has been observed in patients with ARDS (Laurent et al. Am J Respir Crit Care Med 1994;149:1534–8). It is suggested that this may contribute to the excessive pulmonary accumulation of neutrophils observed in ARDS. To assess whether similar mechanisms might be operating in CF, we compared surface expression of CD18 on circulating neutrophils from 10 CF patients (7M, 3F; mean age=20-6, range 21-82 yrs) and 10 control subjects (6M, 4F; mean age=27-7, range 23-35). All CF patients displayed chronic pulmonary bacterial colonisation but none were acutely infected at the time of sampling. As assessed by pulmonary function measurements, lung disease severity ranged from mild (%FEV1 = 85; %FVC = 102) to severe (%FEV1 = 77; %FVC = 44) within the CF study group. Peripheral blood leukocytes were labelled with an FITC-conjugated monoclonal antibody to CD18 and analysed by flow cytometry. Results were calculated as the mean fluorescence intensity (MFI) of CD18-labelled neutrophils. No difference in surface β2-integrin expression on circulating neutrophils between CF patients (MFI = 54-6 (15-8)) and control subjects (MFI = 57-2 (10-8)) was observed. Only one CF patient displayed an MFI value above normal range (i.e. > mean control value + 2SD). Thus, in CF over-expression of β2-integrins on circulating neutrophils does not seem to contribute to the excessive accumulation of neutrophils in the lung. (This work was supported by the Cystic Fibrosis Association of Ireland.)

Effect of recombinant human DNase in vitro and in vivo in bronchiectasis

RA STOCKLEY, VC ORAFOR, C LLEWELLYN-JONES, SL HILL Lung Immunobiological Research Laboratory, The General Hospital, Steelhouse Lane, Birmingham Neutrophil recruitment and the release of elastase into the bronchial secretions is thought to play a significant role in the pathogenesis of chronic bronchiectasis with and without cystic fibrosis. Preliminary clinical studies have investigated the role of inhaled human recombinant DNase in the management of these conditions. However, mucus may entrap several factors that play a part in the pathogenesis of this disease including chemotactic factors and neutrophil elastase itself. Thus disruption of the mucus by DNase may release these factors and adversely affect the lung. We have studied the effect of DNase in vitro and in vivo upon sputum sol phase chemotactic and elastase activity. DNase treatment of five sputum samples in vitro did not affect the amount of albumin recruited or the chemotactic activity; however elastase activity rose from a mean (SE) of 16-2 (3-6) elastase units to 46-3 (8-7) (p<0.05). Twenty bronchiectatic patients entered a double blind study of inhaled DNase (2-5 mg twice a day). Ten received placebo and 10 active treatment. During the run in period the two groups were similar with respect to the degree of lung inflammation as defined by albumin leakage into the lung (mean placebo sputum/serum ratio = 3-35 (1-52) x 10^{-2}; treatment group = 2-35 (0-59)), chemotactic activity (placebo mean = 29-4 (5-4) cells/field; treatment group = 32-3 (2-84)) and elastase activity (placebo mean = 33-7 (7-4); treatment mean = 75-7).
Liposome-mediated CFTR gene transfer to the nasal epithelium of patients with cystic fibrosis

EFPW ALTON, N CAPLEN, PG MIDDLETON, JR DORIN, BJ STEVENSON, X GAO, SR DURHAM, PK JEFFERY, ME HODSON, C COUTELLE, L HUANG, DJ PORTEOUS, R WILLIAMSON, DM GEDDES Ion Transport Unit, National Heart and Lung Institute, London; Department of Biochemistry, St Mary’s Hospital, London; Department of Paediatrics, Papworth Hospital, Papworth, Cambridge; Department of Pharmacology, University of Pittsburgh, USA; Departments of Allergy and Clinical Immunology, Lung Pathology and Cystic Fibrosis, National Heart and Lung Institute, London We have carried out a phase I double blind placebo controlled clinical trial of cystic fibrosis (CF) liposome-mediated CFTR gene transfer in 15 cystic fibrosis (CF) patients. Subjects (n = 3 each group) received either 10 µg, 100 µg, or 300 µg of CFTR cDNA or the appropriate liposome only dose (n = 6), applied topically to each nostril. There were no adverse clinical effects and nasal biopsies demonstrated no histological differences between the groups. Plasmid derived CFTR mRNA was detected in the epithelial layer of the CFTR treated subjects. In vivo assessment of nasal electrophysiological parameters demonstrated an approximately 20% restoration of the deficit (p < 0.05) in both chloride and sodium abnormalities between non-CF and CF subjects. These changes were maximal around day 3 following administration and had reverted to pretreatment values by day 7. In some cases the chloride responses of the CFTR treated subjects reached the range for non-CF subjects. These data are encouraging but suggest that transfection efficiency may need to be improved.

A progressive care programme for patients failing to wean on an ICU: clinical experience

JM SHINEERSON, EM SMITH Respiratory Support Centre, Papworth Hospital, Cambridge During the 32 month period from January 1992-40 patients (21 women), mean (SD) age 54 (16-5) years entered our progressive care programme (PCP) because of a failure to be weaned from mechanical ventilation. The primary diagnosis was pulmonary in 14 patients (COPD 10, post TB sequelae 4). chest wall deformity in five and neuromuscular disease in 21. The mean length of stay on the ICU at time of discharge was 32 days. Twelve patients had failed at least one trial of extubation, 29 patients had been tracheostomised, and 30 patients were completely ventilator dependent. The mean duration of assisted ventilation at the time of transfer was 19-4 hours per day, with 28 patients requiring additional oxygen. Twenty-two patients were treated with non-invasive methods of ventilation and were introduced where possible. The mean length of stay on the PCP was 26-9 (16-2) days. The hospital survival was 95%. Of the survivors 32 patients (84%) were discharged directly to home, three (8%) to residential care, and three (8%) returned to the referring hospital. Twelve patients required no ventilator support at discharge although five retained a tracheostomy. Nineteen patients were discharged with nasal intermittent positive pressure ventilation (NIPPV) and five with negative pressure cuffs ventilation. Of this group of 24 patients five used the equipment for more than eight hours each day. Two other patients remained on IPPV via tracheostomy (one for 16 hours a day and one for 22 hours each day). Only two patients required additional oxygen at home. Mean (SD) overnight SaO2 at discharge was 91 (23-74)% with daytime PaO2 of 10-3 (3-36) kPa and PaCO2 of 6-16 (7-80) kPa breathing air. These findings indicate that most patients who fail to be weaned from mechanical ventilation with conventional techniques can do so and return to their homes with the aid of a comprehensive PCP incorporating the facility of non-invasive home ventilatory support.

Supplementary oxygen administration with nasal intermittent positive pressure ventilation

AJ PADKIN, WM KINNARE Queen’s Medical Centre, University Hospital, Nottingham Intermittent positive pressure ventilation via a nasal mask (nIPPV) can be used to prevent hypercapnia when oxygen is administered to patients who are dependent on hypoxic respiratory drive. We have investigated the inspired oxygen concentration (FIO2) achieved when oxygen is added at different points in the breathing circuit. We studied 11 patients (four women) with chest wall disease, all of whom were accustomed to nIPPV. Oxygen was administered at rates of 1-6 l/min by three different routes: route A, directly into the ventilator (Monnal) with FIO2 remaining constant at a known value throughout inspiration; route B, into the tubing connecting the ventilator to the patient close to the ventilator outlet port; route C, into the nasal mask. By comparing each patient’s SaO2 (Ohmeda 3700) with routes B and C to that obtained with route A we estimated the FIO2 for different oxygen flow rates. The mean FIO2 for route C is shown in the table. There was no significant difference in FIO2 between routes B and C at any flow rate. Route B is preferable to route C in ventilators without the facility for route A since it achieves a similar FIO2 but does not require additional tubing attached to the mask, and is less liable to become disconnected when the patient moves.

Comparison of “elective” and “emergency” initiation of nocturnal assisted ventilation: importance of avoiding endotracheal intubation

PH JOHNSON, BG COOPER, LW L WATSON, JR BRITTON, WM KINNARE Nottingham Assisted Ventilation Group, City and University Hospitals, Nottingham Most patients referred for consideration of nocturnal assisted ventilation (AV) report a gradual decline in health, often over several years. Some of these patients nevertheless present with life threatening acute respiratory failure and require emergency endotracheal intubation. We have compared this group with those presenting earlier, in whom AV can be commenced “electively”. Since 1988 20 patients (eight women) have been commenced on nocturnal AV, 19 with IPPV (18 via a nasal mask and two through a tracheostomy) and one with external negative pressure ventilation. Seven patients required intubation and ventilation on the intensive care unit (ICU) at first presentation. These patients spent a mean (SD) of 21-9 (13-0) days on ICU and a total of 42-9 (11-3) days in hospital. The other 13 patients spent 0-8 (1-5) days on ICU and 11-0 (10-8) days in hospital (p < 0.01). There were no significant differences between the two groups in age, vital capacity, or risk factors for the development of respiratory failure, and the aetiology of respiratory failure was similar. The importance of early referral of patients at risk of nocturnal hyperventilation to an appropriate centre needs to be emphasised to GPs and hospital physicians. At presentation, every effort should be made to use non-invasive AV techniques and avoid endotracheal intubation.

Outcome of nasal intermittent positive pressure ventilation in acute respiratory failure

JS BROWN, DJ MERCHAM JONES, J WEDZICHA Department of Thoracic Medicine, The London Chest Hospital, London Nasal intermittent positive pressure ventilation (NIPPV) has been used in the treatment of acute-on-chronic respiratory failure, although the acceptability and success rate of the technique is variable and has not been evaluated. We report a survey of different hospitals in the UK. Twenty patients treated with NIPPV in 50 patients (25 M, 25 F, median age 69 years, range 45-78) over an 18 month period. Thirty-three patients (66%) had COPD, 11 (22%) restrictive chest wall disease, and six (12%) obstructive sleep apnoea. Thirty-two (64%) patients were using domiciliary LTOT and nine (18%) were on home NIPPV. Twenty-eight (56%) patients had a single admission, 22 (44%) patients had multiple admissions (median 3, range 2-10). Data from the first admission only were included in the analysis. All patients were managed on general wards. Admission (pretreatment) blood
Hypercapnic obstructive acidosis (pH effect of short-term ventilation) HARDINGE, OXFORD. It could well be that failure of their chronic respiratory failure is not statistically significant. Admission pH, PaCO2, and age did not predict failure. Mean (SD) blood gases on NIPPV shortly after admission showed a significant improvement from admission: pH 7.37 (0.07), PaCO2 836 (1.68), PaO2 93 (2.85); mean improvements (95% CI) were pH +0.035 (0.006 to 0.063, p = 0.018), PaCO2 -0.088 (-0.27 to -1.50, p = 0.006), PaO2 +2.51 (1.21 to 3.61, p<0.001). Patients with an admission PaO2 of 6-6 kPa or lower have been identified as at high risk (Jeffrey, Thorax 1992;47:34-40). NIPPV raised PaO2 from an admission level of 6-6 kPa or less in 16 out of 19 patients (p = 0.006). Median duration of NIPPV use was eight days (range 1-47) and of hospital stay 17 days (1-63). NIPPV produced rapid improvement in blood gases in patients treated on general wards for both obstructive and restrictive lung disease. The technique was used successfully in 70% of unselected patients with an acute deterioration of their chronic respiratory failure.

Use of the Hayek oscillator in stable hypercapnic obstructive airways disease FM HARDINGE, JR STARRING. Oster Chest Unit, Churchill Hospital, Oxford. The Hayek oscillator is a new technique which consists of a negative pressure cuirass capable of operating at a variety of high frequencies. It could potentially be used to provide assisted ventilation to patients presenting with respiratory failure resulting from an acute exacerbation of chronic obstructive airways disease. We studied the effect of short term periods of ventilation in 12 subjects with stable hypercapnic chronic airways disease who were outpatients. Subjects received, in a random order, five minute periods of ventilation at three different frequencies, with a “sham” or control period of ventilation at each frequency. Measurements were made before and after ventilation, of changes in gas exchange (oxygen saturation, end tidal CO2), spontaneous respiratory rate (oronasal airflow, inductance plethysmography) and blood pressure (Finapres). In addition, arterial blood gases were measured in four subjects. Each outcome measure was analysed separately for the factors affecting it (ANOVA, Duncan’s post hoc analysis). There was no significant improvement in oxygen saturation, end tidal CO2 or arterial blood gas measurements during any of the “sham” or “active” periods of ventilation. There was significant intersubject variation in respiratory rate (p=0.0001), but no significant changes with ventilation at any setting. No periods of apnoea were observed in any subjects. Blood pressure was also not affected by use of the cuirass. We conclude that the Hayek oscillator does not ventilate patients with stable chronic obstructive airways disease when used for short periods at frequencies of 30-90 oscillations/minute. This is in contrast to experience with normal subjects, and may be due to the decreased chest compliance, altered respiratory drive, and increased anxiety in this group of patients.

Magnetic resonance imaging (MRI) in the staging of bronchial carcinoma AH AL-GHAMDI, CA ROOBOTTOM, M WESTON, C FORBES-WOOD, K JEVINGSHAM, W DHIMIS, P GODDARD, JR CATTLE. Respiratory and Radiology Departments, Bristol Royal Infirmary; Thoracic Surgery Unit and MRI Centre, Frenchay Hospital, Bristol. The role of MRI in the staging of bronchial carcinoma is undefined. Fifty patients considered to have operable bronchial carcinoma on conventional staging including computed tomography (CT) underwent subsequent MRI scanning in the week prior to operation. Axial STIR and T1 weighted images before and after intravenous gadolinium-DPTA and multiecho proton density and T2 weighted coronal images were performed. Assessment of local spread to pleura, chest wall and mediastinum as well as nodal status was made at surgery and by subsequent histopathological examination of resected tissue, and the results compared with the MRI findings. Surgery was performed without the knowledge of the MRI findings. Mediastinal nodes of 0.75 cm or more were demonstrated by MRI with a sensitivity of 92%, specificity of 100%, and accuracy of 96%. Regarding hilar lymphadenopathy the results were 76%, 100% and 85%, respectively. However, lymph node enlargement correlated poorly with the presence of malignancy in the nodes. For pleural involvement the results were 81%, 90% and 87%, respectively. These results are comparable to published results for CT. Regarding mediastinal invasion MRI had a sensitivity of 93%, specificity of 94%, and accuracy of 93%. These results are superior to those reported for CT. In conclusion, MRI is as sensitive as CT in the demonstration of mediastinal and hilar lymph node enlargement, chest wall, and pleural invasion. It is, however, superior in the demonstration of mediastinal invasion and thus holds promise for more accurate staging of bronchial carcinoma preoperatively.

Age as a factor in lung cancer surgery results from one region over a five year period GE WILSON, CJA JACK, ML YLEE, MJ DRAKELEY, RJ DONNELLY, CRK HIND. The Cardiothoracic Centre, Liverpool; Royal Liverpool University Hospital, Liverpool. Lung cancer is now the commonest cause of cancer deaths in men in Great Britain and accounts for 8% of all male and 4% of all female deaths. The prognosis for non-small cell cancer remains poor and surgery remains the best chance of cure. Despite this, only 15% of patients have attempted curative surgery. For a similar stage of lung cancer, patients aged 65 or over are less likely to be operated on. We have therefore looked at the situation over a five year period in Mersey region. A total of 912 thoracotomies (464 lobectomies, 313 pneumonectomies, and 135 wedge resections) were performed from August 1987 to July 1992. We examined the records of all patients aged 70 or above undergoing surgery for lung cancer (elderly). A random group of patients below this age was selected for comparison (young). Seventy nine elderly patients (50 men; mean age 75, range 70-87) and 83 young patients (53 men; mean age 60, range 34-69) underwent thoracotomy for bronchial carcinoma. The characteristics for the two groups are as shown in the table. The type of operation performed, tumour cell type and stage were similar for each group. Postoperative complications were fewer in the elderly (26%) than in the young (38%), due mainly to chest infections. Adjuvant radiotherapy was used in 26% of elderly and 18% of young patients. The mean survival for each group was 23 (2) months. From these figures it would appear that age should not necessarily be considered a contraindication to surgery for lung cancer. Furthermore, patients over the age of 70 have fewer postoperative complications and equal survival figures to younger patients, despite having to wait longer for referral.

Does blood transfusion affect long term survival following lung cancer resection? RK MACKENZIE, MM KENNEDY, M GREISS, KM KERR, RR JEFFREY. Departments of Cardiothoracic Surgery, Transfusion Medicine and Pathology, Aberdeen Royal Infirmary NHS Trust, Aberdeen. Debate continues on the potential deleterious effects of perioperative blood transfusion in patients undergoing potentially curative resections for cancer. We have studied the results of 280 patients who underwent pulmonary resection for non-small cell lung cancer between January 1986 and January 1991 looking at the effect of perioperative blood transfusion on long term survival. Demographic data were obtained from the case records, perioperative transfusion data were obtained from the Regional Blood Transfusion Service records, and the pathology was reviewed. Date of death was obtained from the Scottish Cancer Registry. Patients dying within 90 days of operation (n = 26) were excluded. The mean age of the group was 62.3 (8.8) years and 65.7% were men. There were 116 squamous carcinomas, 89
adenocarcinoma, 21 of mixed differentiation, 11 large cell carcinomas, and 17 others. Eighty-five patients underwent pneumonectomy, 148 lobectomy or segmentectomy procedures, e.g., segmentectomy. Three year survival for pneumonectomy was 44.7% compared with 60.5% for lobectomy; 125 (49.2%) patients had stage I disease, 79 (31.1%) stage II and 50 (19.7%) had stage III. Patients with stage I disease survived a mean of 46.4 (24.2) months with a three year survival of 60.8%, those with stage II 40.0 (25.8) months and 53.8%, and for stage III 31.1 (26.4) months and 40%. Over all stages patients with adenocarcinoma did marginally worse than those with squamous tumours (e.g. stage I 67.3% vs 63.8%). For all stages of disease patients transplanted perioperatively had significantly reduced long term survival e.g. stage I not transplanted 58.6 (26.3) months v transplanted 40.2 (4.4) months, p<0.001. We conclude that perioperative blood transfusion should be considered among the major determinants of long term survival following resection for lung cancer.

Patient survival and bronchial resection line status in primary lung carcinoma

KK TAN, MM KENNEDY, KM KERR, RR JEFFREY Departments of Cardiothoracic Surgery and Pathology, Aberdeen Royal Infirmary A series of 255 patients who had lung resection for primary non-small cell cancer between 1986 and early 1991 were studied. Postoperative survival was calculated from Cancer Registry data; patients surviving <3 months were excluded. All histological material was reviewed. From 255 cases 18 (7%) showed tumour at the bronchial resection line (BRL), six (2.4%) cases showed carcinoma in situ (CIS), and five (2%) showed dysplasia. Of the 18 cases of BRL with tumour five showed mucosal disease only while 13 showed extrachondral tumour. Mean survival for those with invasive disease at the BRL was 32.6 (24-7) months compared with 41.7 (26-0) months for those without (p=0.0004). There was no significant difference in mean survival between the groups with CIS, dysplasia, or benign BRL, respectively.

The case group was one mucosal group at the BRL but all the groups with extrachondral tumour at this site, had 40% of patients alive at the review date (>46 months postoperatively). Of those who died before the review date mean postoperative survival with only mucosal disease at BRL was 29.1 months, while it was only 11 months for the group with BRL showing extrachondral cancer. These data suggest that invasive disease, and particularly extrachondral malignancy, but not carcinoma in situ or dysplasia may be a relevant factor in determining patient survival after lung resection for primary bronchial carcinoma.

TNM staging system for malignant mesothelioma: evaluation in 86 patients with malignant pleural mesothelioma

L TAMMILEHTO, K MATTSON, L KIVISAAR, U-S SALMENP, P MAASILTA Department of Epidemiology and Biostatistics, Institute of Occupational Health, Helsinki; Departments of Pulmonary Medicine, Diagnostic Radiology, and Thoracic and Cardiovascular Surgery, Helsinki University Central Hospital, Helsinki, Finland There is no universally recognised method for staging malignant mesothelioma, although computed tomographic (CT) scanning has improved non-invasive staging. IUAC has recently proposed using the TNM staging system for mesothelioma, but in clinical practice it is difficult to assess the tumour (T) and nodal (N) characteristics of pulmonary mesothelioma. The staging growth pattern of this tumour. In order to evaluate TNM staging we analysed preoperative CT scans from 86 patients with histologically confirmed malignant pleural mesothelioma. Most of the patients would participate in a clinical study programme which included debulking surgery, chemotherapy and hemithorax irradiation. Median age of patients was 56 years (range 38–79). There were 70 men and 18 women, and 33 had tumours with epithelial histology. Median survival for all the patients, measured from the data of histological confirmation of mesothelioma, was 10 months (range 0.2–110). The same radiologist (LK) analysed CT scans according to the TNM staging system. Actuarial survival curves were constructed by the Kaplan-Meier method. Survival curves for the different TNM categories were compared using the log rank test. Node evaluation could not be completed in eight cases because the tumour had encompassed the hilum and mediastinum. In multivariate analysis significant differences in prognosis correlated to the different T categories (p<0.01), and the different stage categories (p<0.05), but not to the N or M categories. Larger studies are needed to assess the importance of TNM staging for selecting treatment and as a prognostic factor for malignant mesothelioma.

Eight year experience with the pleuroperitoneal shunt for the management of malignant pleural effusions

M PETROU, D KAPLAN, P GOLDSTRAW Royal Brompton Hospital, London Malignant pleural effusions are a common cause of morbidity and patients often require repeated palliation by needle thoracocentesis, tube thoracostomy, and pleurethorax. For patients where lung expansion is restricted by a fibrous or malignant cortex ("trapped lung") pleurethorax usually fails and therefore a shunt is needed. Over an eight year period (1986–94) 275 patients were referred to our surgical unit with pleural malignancies, 65 of whom had no significant effusion (underwent pleural biopsy only) or were looking to have an empyema (requiring rib resection and external drainage). This left 210 patients with large recurrent effusions and 128 were treated by talc pleurethorax. We review the results of the remaining 82 patients who were treated with a pleuroperitoneal (Denver Biomedical Inc) shunt for recurrent malignant effusions associated with a trapped lung. The mean age was 60 years (range 25–85 years) and 62 patients (76%) had received repeated first line treatments before referral. The primary tumours were mesothelioma (39 patients), secondary breast carcinoma (15 patients), and primary or secondary adenocarcinoma of the lung (12). All patients underwent general anaesthesia and thoracotomy or mini-thoracotomy. The degree of lung expansion was assessed by sustained positive pressure ventilation and the patients were eligible to be subjected to a pleuroperitoneal shunt in all patients resulting in the insertion of a pleuroperitoneal shunt. There were no intraoperative deaths and only two early deaths (2.4%).

The mean hospital stay was 5.9 days (range 2–12 days). Follow up data were complete in 85% of the patients. There were nine cases (4.5% of all shunts) requiring replacement or renovation and four requiring removal for sepsis) at one week to four months after insertion. One patient required further thoracocentesis and two patients developed malignant ascites. The median survival in this heterogeneous group of cancer patients was 5 months (range 1–36) and those with secondary breast carcinoma or lymphoma surviving the longest. Our results show that patients in whom pleurethorax would fail due to limited lung expansion despite adequate drainage of the effusion can obtain safe and effective palliation by pleuroperitoneal shunting.

Tetracycline pleurethorax in malignant pleural effusion: a comparison of needle aspiration with intercostal tube drainage

LG MICALPINE, JW KAY, NC THOMSON, BHR STACK, PA CORRES, E NEVILLE, FJC MILLARD ON BEHALF OF THE BRITISH THORACIC SOCIETY RESEARCH COMMITTEE The British Thoracic Society, St Andrews Place, London Chemical pleurethorax is often required to prevent re-accumulation of malignant pleural effusion. Tetracycline is the commonest agent used but the method of removing the pleural fluid varies (Micalpine et al, Thorax 1990;45:699–701). The present study compared the success of tetracycline pleurethorax using a needle aspiration technique with an intercostal tube drainage method in patients with recurrent malignant effusion. Patients with recurrent pleural effusion due to histologically or cytologically proven malignancy other than mesothelioma or lymphoma who were suitable for pleurethorax (lung volume reduction) were eligible for this multicentre trial. They were randomised to either needle aspiration of the effusion to dryness followed by instillation of 1.5 g tetracycline in 50 ml saline containing 200 mg lignocaine or to intercostal tube drainage to dryness prior to instillation of the same mixture and drainage then continued for >48 hours or until <100 ml/day. A total of 113 patients were randomised but 32 were lost to follow up or were withdrawn. Thus there were 81 evaluable patients (lung cancer 34; breast 15; other/unknown primary 32). Forty one were allocated needle aspiration and 40 an intercostal tube drainage. Pleurethorax was successful (no need for further aspiration) at six weeks in 11 of 35 (32%) in the needle aspiration group and in 29 of 36 (80%) in the intercostal tube drainage group (p<0.0001). At 12 weeks the corresponding success rates were seven of 33 (21%) and 25 of 32 (78%) (p<0.0001), and at 24 weeks four of 30 (13%) and 19 of 27 (70%) (p<0.0001). There were five failures in each treatment group due to problems with the technique. The underlying diagnosis did not influence success. There were no differences in the time between pleurethorax and hospital discharge,
in the complication rate, or in the acceptability to patients between the two techniques. In conclusion, intercostal tube drainage of pleural fluid when carrying out tetracycline pleurodesis for malignant effusion is considerably less invasive than the pleural needle aspiration method without being any less acceptable to patients.

Mast cells infiltrate the bronchial epithelium in mild asthma

M SYNEK, AJ FREW, FC LAMPE, P BRADDING, ST HOLGATE University of Southampton, Southampton We compared numbers and distribution of mast cells infiltrating the airways in eight mild asthmatic subjects who died of causes unrelated to asthma with nine non-asthmatic subjects who died of incidental causes. Specimens of lung tissue were obtained at post-mortem examination. Sections 4 μm thick were cut from the tissue blocks and stained by monoclonal antibody (AA1:tryptase Moab) using streptavidin-biotin peroxidase technique. The numbers of positive cells were expressed in cells/mm² for the epithelium and cells/mm² for the submucosa. The airways were divided into two groups according to the internal perimeter (P): those with P<1.5 mm and those with P>2.5 mm. All airways together were studied first, then larger and smaller airways were evaluated separately. When all airways were examined together we found higher numbers of mast cells infiltrating the epithilium in mild asthma than in non-asthmatic subjects (2-06 ± 0.17 cells/mm², p = 0.006). In the submucosa the numbers of mast cells were not significantly different (103-4 ± 112-7 cells/mm², p = 0.597). The same applied to larger and smaller airways examined separately. The respective figures for epithelium were 1-31 ± 0.23 cells/mm², p = 0.03, and 1-99 ± 0.03 cells/mm², p = 0.02. The corresponding figures for the submucosa were 103-2 ± 106-1 cells/mm², p = 0.80 and 130-6 ± 159-0 cells/mm², p = 0.42. These results indicate that mast cells infiltrate the bronchial epithelium in mild asthma, while the numbers of cells in the submucosa do not differ from normal subjects. This distribution of mast cells is present throughout the bronchial tree, irrespective of airway size.

Allergen-induced late asthmatic reactions are associated with increased nitric oxide concentration in exhaled air

SA KHARTONOY, DJ EVANS, BJ O'CONNOR, PJ BARNES Department of Thoracic Medicine, National Heart and Lung Institute, London; Clinical Units, St George's Hospital and Chelsea and Westminster Hospital. The concentration of nitric oxide (NO) is increased in the exhaled air of asthmatic patients and may reflect cytokine-mediated inflammation in the airways. We have used inhaled allergen challenge (AC) to induce a late asthmatic reaction (LAR) to explore whether allergen-induced inflammation causes an elevation of NO in asthma. We used serial FEV1, by spirometry and NO using a chemiluminescence analyser for up to 27 hours following allergen or methacholine challenge and measured responsiveness to inhaled histamine by determining the provocative concentration causing a 20% fall in FEV1 (PC20) before and three hours after challenge. Of 22 patients who underwent AC, 16 developed dual early and late responses, whereas six had a single early response. In the patients with a dual response the mean maximal fall in FEV1, during the LAR was 34-9 (8-6%) at 21 hours after challenge and there was a significant reduction in histamine PC20 at three hours. In these patients there was a significant increase in the level of NO, with a mean increase of 1.8 fold (range 1.7-10.5) hours after challenge. There was a significant relationship between the size of the LAR and increase in NO (r = 0.75, p<0.01). In patients who had a single early response, which was not significantly different from the early response in dual responders, there was no significant change in NO over the 27 hour period or in histamine PC20. In five patients given a control challenge with methacholine rather than AC and followed over the same period there was no change in NO. Similarly, there was no increase in NO after inhaled histamine in any of the patient groups. We conclude that patients who develop a LAR after AC show an increase in the amount of NO in exhaled air which precedes the fall in lung function, whereas patients who have an isolated early response to allergen do not show any change in NO. Similarly challenge with saline or histamine has no effect on exhaled NO. This provides further evidence that exhaled NO may reflect allergic inflammation in asthmatic airways, and may be a useful marker in monitoring asthma and its response to anti-inflammatory treatment.

Imaging allergen-invoked airways inflammation in atopic asthma in vivo using [15F]-fluoroethoxyglucose and positron emission tomography (PET)

IK TAYLOR, AA HILL, M HAYES, C RHOADES, T JONES, JBM HUGHES, NB PRIDE, RW FULLER Departments of Respiratory Medicine, Clinical Pharmacology and MRC Cyclotron Unit, RPHMS, Hammersmith Hospital; Glaxo Group Research Airways inflammation is a cardinal feature of chronic asthma but is currently only visualisable by bronchial lavage and biopsy. Inflammatory foci can, however, be imaged non-invasively using PET and [15F]-labelled fluoroethoxyglucose ([15F]FDG, a non-metabolisable analogue of D-glucose) to quantify glucose uptake of activated granulocytes. We have investigated the effect of segmental allergen challenge on FDG uptake in the lungs of five mild male asthmatic patients (age 23-32, FEV1, 71-110% predicted, PC20 histamine (H) < 16 mg/ml, on inhaled β2 agonists alone). On day 1, pre-allergen PC20 H was measured. The following day at bronchoscopy 20 ml allergen (D. pteronyssinus or mixed grass pollen, range 100-500 units, median 200) was instilled into the posterior segment of the right upper lobe (RUL, time 0). A similar volume of saline was instilled into the posterior segment of the LUL. At variable times following allergen instillation PET scanning was performed with the patients resting supine within the scanner. Following an initial transmission scan, about 3 mCi of [15F]FDG was injected intravenously. Blood was sampled from the antecubital vein just before the uptake scan and sequential time frames of [15F]FDG radioactivity (each of 15 x 5 mm planes) were acquired over the next 66 minutes. Regions of interest (to permit calculation of the rate of uptake of [15F]FDG) were defined anatomically and were drawn manually from the initial transmission scan to incorporate the posterior two thirds of the lung fields from the first four planes above the main carina. Rate of uptake of [15F]FDG (min^-1) was calculated from Patlak plots in which tissue/plasma count ratio (x 10^-3) was plotted against the integral plasma counts/plasma counts to obtain a slope; the RUL/LUL (R/L) slope ratios were then calculated. Following completion of the PET scan, post allergen PC20 H was measured and the change in airway reactivity expressed as the doubling dose (DD) shift. [15F]FDG uptake was significantly increased in all studies in the RUL compared to the LUL: slope RUL 6-1-26-7 x 10^-3 min^-1 median 12-0 ± 3-80; median 2-6, R/L slope ratios 1-3-22, median 4-0, p<0.02. Median DD shift was 3-0-76%. No local allergen-induced changes could then be visualised using [15F]FDG and PET in vivo in asthma. The cellular localisation of the [15F]FDG signal however remains to be determined.

Human bronchial epithelial cell mediators influence eosinophil and neutrophil adhesion to human endothelial cells in vitro

MM ABEDELZAEZ, JL DEVALIA, OA KHARIJ, PJ SAPPINORD, PJ DAVIES Department of Respiratory Medicine and Allergy, St Bartholomew's Hospital, London; Department of Thoracic Medicine, National Heart and Lung Institute, London; Glaxo Group Research Airways inflammation is a cardinal feature of chronic asthma but is currently only visualisable by bronchial lavage and biopsy. Inflammatory foci can, however, be imaged non-invasively using PET and [15F]-labelled fluoroethoxyglucose ([15F]FDG, a non-metabolisable analogue of D-glucose) to quantify glucose uptake of activated granulocytes. We have investigated the effect of segmental allergen challenge on FDG uptake in the lungs of five mild male asthmatic patients (age 23-32, FEV1, 71-110% predicted, PC20 histamine (H) < 16 mg/ml, on inhaled β2 agonists alone). On day 1, pre-allergen PC20 H was measured. The following day at bronchoscopy 20 ml allergen (D. pteronyssinus or mixed grass pollen, range 100-500 units, median 200) was instilled into the posterior segment of the right upper lobe (RUL, time 0). A similar volume of saline was instilled into the posterior segment of the LUL. At variable times following allergen instillation PET scanning was performed with the patients resting supine within the scanner. Following an initial transmission scan, about 3 mCi of [15F]FDG was injected intravenously. Blood was sampled from the antecubital vein just before the uptake scan and sequential time frames of [15F]FDG radioactivity (each of 15 x 5 mm planes) were acquired over the next 66 minutes. Regions of interest (to permit calculation of the rate of uptake of [15F]FDG) were defined anatomically and were drawn manually from the initial transmission scan to incorporate the posterior two thirds of the lung fields from the first four planes above the main carina. Rate of uptake of [15F]FDG (min^-1) was calculated from Patlak plots in which tissue/plasma count ratio (x 10^-3) was plotted against the integral plasma counts/plasma counts to obtain a slope; the RUL/LUL (R/L) slope ratios were then calculated. Following completion of the PET scan, post allergen PC20 H was measured and the change in airway reactivity expressed as the doubling dose (DD) shift. [15F]FDG uptake was significantly increased in all studies in the RUL compared to the LUL: slope RUL 6-1-26-7 x 10^-3 min^-1 median 12-0 ± 3-80; median 2-6, R/L slope ratios 1-3-22, median 4-0, p<0.02. Median DD shift was 3-0-76%. No local allergen-induced changes could then be visualised using [15F]FDG and PET in vivo in asthma. The cellular localisation of the [15F]FDG signal however remains to be determined.
ICAM-1 expression on alveolar macrophages (AM) is enhanced by relevant allergen challenge in patients with atopic asthma

M SARNO, S SETHI, J ALLEN, S ROTONDETO, R KNIGHT, M SPITERI Lung Cell Biology Group, Department of Postgraduate Medicine, Keele University, Staffordshire The intercellular adhesion molecule 1 (ICAM-1) has been postulated to play a critical role in the intricate cell–cell interactions between immune cells and epithelial cells which line the airways in patients with atopic asthma. In this study, surface ICAM-1 expression was analysed using a monoclonal antibody RB1/1-1 (Boehringer Ingelheim, USA) in freshly isolated unstimulated and allergen-stimulated bronchoalveolar lavage cells (BALC) from six atopic asthmatic and six allergic asthmatic patients. A control group of six healthy volunteers was included. Cultures were set up in serum-free RPMI 1640 medium in 35 mm dishes (1 x 10⁶ cells/dish) for 24 hours. The immunostaining intensity was assessed using a semiquantitative visual score from two counts of 300 cells per slide. Baseline expression of surface ICAM-1 (10 ng/ml) was preferentially increased in a concentration-dependent manner from a control of 83 (3)% viability with GM-CSF alone to 39 (10)% with CTX (1 μg/ml) (CV = 7 mg/ml) (n = 4). The adenylate cyclase activator, forskolin, however, did not inhibit ICAM-1 expression at any concentration tested (10⁻¹⁻⁻ to 10⁻⁵ M). The expression of ICAM-1 on GM-CSF-stimulated BALC was dependent on the presence of cyclic AMP (cAMP) that elevate intracellular cAMP levels (1 mM forskolin, (30) = 200 μM). It was found that GM-CSF increases ICAM-1 expression on AM preferentially in a concentration-dependent manner from a control of 59 (12) fmol/10⁶ EOS to 515 (98) fmol/10⁶ EOS with CTX (1 μg/ml) (n = 3). Forskolin, however, only increased CAMP at the highest concentration tested (10 μM) from a control of 69 (20) fmol/10⁶ EOS with GM-CSF alone to 181 (30) fmol/10⁶ EOS with forskolin (n = 5). Two cell lines of cytokines undergo apoptosis, which damage characteristic of their DNA. We have examined the DNA from cells cultured for seven days and found that GM-CSF (1 μg/ml) prevents this DNA breakdown and the protective effect of GM-CSF can be overcome with the induction of DNA fragmentation by agents that elevate intracellular CAMP (1 mM dibutyryl cAMP and 10 ng/ml CTX).

ICAM-1 expression on alveolar macrophages (AM) is enhanced by relevant allergen challenge in patients with atopic asthma

C O'LEARY, J DANIELS, PW JONES FOR THE ISOLDE GROUP Division of Physiological Medicine, St. George's Hospital Medical School, London; Department of Respiratory Physiology, Birmingham Heartlands Hospital, Birmingham The relationship between spirometric parameters and impaired quality of life in COPD is weak, but the size of the ISOLDE study (concerning the effect of fluticasone propionate on long-term outcome in COPD) allows more detailed analysis than was possible previously. Spirometry was performed using a computer-controlled rolling seal device (Sensormedics) to ATS quality criteria. The mouthpiece was attached by a "no-flow" point (i.e. FVC). Spirometry was performed four hours after any previous bronchodilator, then following 400 μg salbutamol plus 80 μg ipratropium delivered via a Nebuhaler (10 breaths). Baseline data from 273 patients (71 women) were analysed. Their mean (SD) age was 64 (4) years, mean postbronchodilator FEV₁ was 51.1 (13.9) % pred, and total St George's Respiratory Questionnaire (SGRQ) score was 51.18. Change in FEV₁ following bronchodilators (ΔFEV₁) was 6.7 (5.3)% pred. Pearson correlations between SGRQ and spirometric values are shown in the table. The correlation between SGRQ score and ΔFEV₁,

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<td>FVC (% pred)</td>
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(As % pred) was significant (r = -0.18, p < 0.05). In a multiple regression with score as the dependent variable both FEV₁ and FVC were significant (F = 7.4), but not FVC. In the absence of any bronchodilator treatment, the SGRQ score correlated better with the best FEV₁, rather than the worst – probably a result of the standardising effect of bronchodilator. The absence of any correlation between SGRQ and FVC was surprising and may have been due to the use of strict criteria for values, and the "no-flow" point to determine the "no-flow" point prior to each FVC. Thus, this suggested that patients with severe breathlessness had no long-term benefit of bronchodilator treatment, and thus increased oxygen consumption and therefore increased respiratory effort may not influence respiratory function in patients with COPD.

Treatment of breathlessness without incurring sedation or respiratory depression in anxious patients with COPD using Buspirone

AK DATTA, S MOUZENI, N GENTILE, S MCAFEE, K MACMURRIS, P JONES, A MACRAE. Department of Medicine, Charing Cross & Westminster Medical School, London; St George's Hospital Medical School, London; Clinical Professor of Medicine, University of London; Consultant Respiratory Physician, Brompton Hospital; Senior Lecturer in Medicine, University of Birmingham. The development of a new non-addictive drug to treat anxious patients with COPD is needed. This was a three-centre, double-blind, placebo-controlled, randomised, crossover study. After informed consent was obtained, 28 patients were randomised and each received 200 mg buspirone or placebo in a 1/1 ratio. Both treatments were administered during exercise (90 watts for 4 minutes) to a respiratory task. The patients were monitored breath by breath during exercise using a breath-by-breath computer system. After each exercise, the patients were questioned about their subjective experience of the exercise. After the exercise period, the patients were then asked to rate their anxiety, dyspnoea and chest discomfort during the exercise using visual analogue scales. The results showed that buspirone reduced anxiety and did not affect exercise performance, whereas placebo increased anxiety and decreased exercise performance. These results suggest that buspirone is a potential new treatment for anxious patients with COPD. Further studies are needed to confirm these findings.
ventilation at rest or when stimulated by hypercapnia or exercise. There is a modest but statistically insignificant reduction in perceived breathlessness at rest. During exercise patients appeared to tolerate a greater level of metabolic work and exercise related oxygen desaturation.

Magnetic stimulation of the diaphragm in patients with chronic hyperinflation

MI POLKEY, D KEROYSSUS, SJ KEELY, GH MILLS, CH HAMMENGERD, M GREEN, J MOKHAM King's College Hospital and Royal Brompton Hospital, London Patients with COPD have hyperinflation and shortening of the diaphragm. This should theoretically result in reduced force and pressure generating capacity, but other investigators (Similowski et al. N Engl J Med 1991;325:917–23) have found evidence of better than expected contractility suggesting a process of chronic adaptation. We therefore used cervical magnetic stimulation of the phrenic nerve roots to study the effect of chronic hyperinflation on diaphragm contractility in seven well trained men with severe COPD (mean FEV1, 0·64 l, range 0·5–0·9). Lung volumes were determined using whole body plethysmography. To avoid diaphragm potentiation (Am J Respir Crit Care Med 1994;149:739–43) the subjects breathed quietly for 20 minutes after passage of oesophageal and gastric balloon catheters. Twitch transdiaphragmatic pressure (Tw Pdi) was then measured at resting and expiration. Five measurements were made for each subject. Tw Pdi was reproducible for each subject (coefficient of variation 2·7–7·5%). The mean Tw Pdi was 20·3 cm. H2O (range 14·9–25·2) at a mean lung volume of 2·1 litres (range 0·9–3·2) above predicted FRC. In normal male subjects in our laboratory (Am J Respir Crit Care Med 1994;149:A133) the mean Tw Pdi at FRC was 31·8 cm. H2O (range 26·5–38·1), but at 2·1 litres above FRC the mean Tw Pdi was 20·5 cm. H2O. We conclude that during chronic hyperinflation diaphragm contractility is comparable to that seen in normal subjects during acute hyperinflation and we observed no evidence of chronic adaptation.

Measurement of maximal exercise capacity in patients with severe chronic obstructive pulmonary disease (COPD)

RS MATHUR, SM REVILL, DD VARA, RW WALTON, MDJ MORGAN Department of Respiratory Medicine, Glenfield Hospital, Leicester In normal subjects treadmill exercise usually produces the greatest maximal oxygen consumption (VO2 max). This may not be true in COPD where bicycle exercise, which supports the upper limb muscles, may be better tolerated than unsupported walking on a treadmill. The aim of this study was to determine which mode of exercise produced the best performance in patients with severe COPD. Eight patients with severe COPD (mean FEV1, 28% predicted, range 18–36) exercised to a symptom limited maximum on a cycle and on a treadmill on separate days. Cycle workload was increased by 10 watts/minute and the treadmill gradient was increased by 2·5% alternate minutes, the speed remained constant. VO2, ventilation (Ve), and heart rate (HR) were measured. Lactate concentration was measured at rest and four minutes after exercise. Breathlessness (BS) and perceived exertion (PE) were rated from modified Borg scores. Full pulmonary function tests were carried out during a screening visit. The mean (SE) VO2, Ve and HR at peak cycle exercise were 11·8 (1·1) ml/min/kg, 27·3 (3·2) l/min and 120 (5·5) b/min respectively, and for peak treadmill exercise 12·0 (1·1), 26·0 (2·8), and 121 (7·4). There were no significant differences, though individual variability was large. There was a difference between the exercise lactate concentrations, with cycle exercise generating the greatest response 2·49 (0·42) vs 1·22 (0·27) mmol/l, (p<0·05). There was a better correlation between the Borg scores and VO2 during cycle exercise than during treadmill exercise (0·700 for BS and 0·784 for PE v 0·448 and 0·495). The relationships between static lung function and exercise performance were poor. There was no clear difference between the two forms of exercise. Individual variation was large therefore the best performance cannot be predicted. This can only be determined if both types of exercise are carried out. Cycle exercise was unfamiliar to the patients and generated the greatest lactate response. The results do not support the hypothesis that cycle exercise consistently produces a better performance in severe COPD. [RSM sponsored by Raj Nanda Pulmonary Disease Research Trust Delhi, and The British Thoracic Society.]

Metabolic and cardiorespiratory responses to inspiratory threshold loading in COPD

SM REVILL, MDJ MORGAN, AE HARDMAN Department of Respiratory Medicine, Glenfield Hospital, Leicester; Department of Sports Science, Loughborough University The aim of the study was to examine the metabolic and cardiorespiratory responses to inspiratory threshold loading (ITL) in patients with COPD. A solenoid device was used to generate the threshold pressure. Its use has been described in normal subjects (Bardsley PA et al. Thorax 1993;48:354–9). Eleven patients were recruited (8M, mean age 61) with moderate COPD (mean (SD) FEV1, 1·29 (0·43)). All patients carried out a practice test and seven carried out an additional test to determine repeatability. Patients breathed through a mouthpiece attached to the solenoid device and the gas analysis system. VO2, Ve and heart rate (HR) were measured at rest, and with mouth pressure (Pm) and the pressure time index (Pdi) during ITL. The initial load and size of increments were determined for each patient, aiming to achieve a maximum within 10 minutes. The load was increased every minute until the patient could no longer inspire. A blood sample was taken at rest and four minutes after loading to determine lactate concentration. The repeatability of peak Pm and Pdi at the end of the test was good but the VO2 peak and Ve peak were more variable. The mean difference and 95% limits of agreement (LA) for tests 2 and 3 are shown in the table. For the whole group (n=11) the mean (SE) peak Pm was 52·7 (7·0) cm. H2O, Pdi 1410 (261) cm. H2O/s, VO2 peak 367 (17) ml/min, Ve peak 24·2 (1·9) l/min, and HR peak 89 (5·7). The mean (SD) increase from resting levels for VO2, Ve and HR were 118 (52) ml/min, 11·4 (6·5) l/min and 12·4 (6·0) (p=0·05). These represent mean percentage increases of 47, 96, and 16, respectively. There was no significant difference in lactate concentration. Threshold loading in COPD using the solenoid device was reproducible. The incremental, progressive protocol produced significant increases in VO2 peak, Ve peak, and HR, but no lactate response.

Bicycle ergometer underestimates symptom-limited maximal VO2 in subjects with moderate chronic obstructive pulmonary disease (COPD): implications for clinical testing and rehabilitation programmes

AN MCLEAN, LM COCHRANE, CJ CLARK Hairmyres Hospital, East Kilbride, Lanarkshire Cycling and walking are used as means of assessing exercise capacity and exertional dyspnoea and to improve exercise tolerance in pulmonary rehabilitation programmes. The choice of exercise mode is often arbitrary. The subjective differences in the patient's and investigator's physiological responses between the two modes are subjects with COPD are not known. We examined the effect of each mode of exercise on the relationships of: (1) breathlessness (Borg rating) with oxygen uptake (VO2); (2) minute ventilation (Ve) with VO2; and (3) Ve with Borg. We also assessed symptom limited maximal VO2 between modes in subjects with COPD. Nine subjects with moderate COPD (mean FEV1, 45% pred) performed a symptom-limited progressive incremental test on bicycle (B) and treadmill (TM) on the same day. Regression analysis was carried out for each subject for each relationship on both TM and B. The mean slope and intercept between the modes was compared using the paired t test. For Borg vs VO2 there was no significant difference, slope TM 0·0098 ± 0·0088, constant TM −6·25 ± 5·36. For Ve vs VO2 Ve was lower on the TM, slope 0·026 ± 0·030 (p<0·01), constant 3·88 ± 3·30. For Borg vs Ve, Borg was higher on TM, slope 0·328 ± 0·274 (p<0·01), constant −5·28 ± 6·75. Maximal VO2 was higher on TM, 1·53 ± 1·29 (p<0·001) but there was no significant difference in maximal Ve (40·2 ± 44·1 l/min). Subjects were working close to their predicted maximal voluntary ventilation in both modes with dyspnoea indices of 89% ± 91% for B and TM respectively (NS). This study shows that less work was achievable on the bike which may significantly underestimate symptom-limited maximal VO2 in these patients. This has not been previously described in this group of subjects in whom
the respiratory limitation might be expected to predominate and prevent differences in maximal work capacity from appearing. The submaximal data show a difference in ventilatory response between the two modes but suggest that subjective breathlessness remains closely linked to VO2 irrespective of the mode of exercise.

Outcomes in occupational asthma

DJ ROSS, JC MCDONALD SWORD Project, National Heart and Lung Institute, London SWORD (surveillance of work-related and occupational respiratory diseases) is a voluntary reporting scheme in which 83% of consultant chest physicians in the UK participate as well as a similar number of occupational physicians. We have undertaken a study of the long term respiratory health and employment after a diagnosis of occupational asthma. A total of 1940 cases of occupational asthma in the age group 16–65 were notified to the scheme from 1989–1992. A questionnaire was sent for each case to the 312 physicians who reported them. Duplicates, cases where the physician had retired, or the diagnosis had been revised were excluded. So far, 81% of questionnaires have been returned, 792 by chest physicians (35% initially medicolegal) and 231 by occupational physicians (4% medicolegal). Time from notification to response was 17–68 months. 750 cases were not medicolegal. Of these, 21% were no longer considered to have asthma, 74% had improved, 22% were unchanged, and 4% were worse; 48% were with the same employer and 13% were employed elsewhere, 23% were unemployed, 7% medically retired or not working and 4% retired. Unskilled workers had a higher probability of leaving their employer than skilled workers. The mean duration of exposure to the suspected agent was 20–0 months in those who recovered from asthma, but was 52–6 months in those still affected. The mean duration of exposure after diagnosis was 5–7 months in those who recovered, but 11–0 months otherwise.

Work-related symptoms and specific sensitisation in a cohort of workers exposed to acid anhydrides

RD BARKER, M VAN TONGEREN, JM HARRIS, KM VENABLES, AJ NEWMAN TAYLOR Department of Occupational and Environmental Medicine, National Heart and Lung Institute, London; Institute of Occupational Health, Birmingham To study exposure response relationships in occupational asthma we have examined a cohort of workers exposed to acid anhydrides (AA) at four industrial sites in the UK. Participants completed questionnaires on respiratory symptoms, smoking habits and work history and had skin prick tests (SPT) with common inhalant allergens and three AA-human serum albumin (AA-HSA) conjugates. In 1992 exposure to phthalic anhydride (PA), trimellitic anhydride (TMA), and maleic anhydride (MA) was measured with personal samplers during full shifts. Past exposure was estimated using job and process descriptions and past exposure measurements. The risk of being sensitised or experiencing new work-related chest symptoms was assessed in relation to full shift AA exposure. In total, 506 workers fulfilled the cohort definition of whom 401 (79%) took part (388 (97%) men; age range 18–81, median 39 years). Thirty-four (9%) had new work-related chest symptoms and 12 (3%) had a positive SPT to an AA-HSA conjugate. Average full shift exposure to all AA, measured in 1992, was below the occupational exposure limit at all sites. There was a trend for an increasing prevalence of sensitisation with increasing exposure to TMA (z-test for linear trend p=0.019). This trend was not apparent when all 32 AA were included in the analysis. The highest average full shift exposure experienced by five workers sensitised to TMA was less than one quarter of the current UK occupational exposure limit. A matched case–control analysis showed an increased risk of new work-related chest symptoms with increasing estimated full shift exposure to acid anhydrides. The data show that higher full shift exposures are associated with a greater risk of sensitisation and symptoms. They also suggest that the current occupational exposure limit for TMA may be too high.

Coal miners aged over 70 seeking industrial injury benefit for bronchitis and emphysema

PS SANDHU, SC STENTON, SJ BOURKE, DJ HENDRICK Chest Unit and Regional Unit for Occupational Lung Disease, Newcastle General Hospital, University of Newcastle upon Tyne; Benefits Agency Medical Service Industrial injury benefit for coal miners suffering from chronic bronchitis and emphysema was introduced in September 1993. Miners with at least 20 years underground exposure, an FEV1 of at least 1 litre below the Cotes predicted values, and pneumoconiosis (ILO category 1/0 or above) are eligible for benefit. To date, over 7300 miners have been evaluated at the Newcastle office of the Benefits Agency. We have collected data on respiratory symptoms, exposure and smoking histories, lung function and radiographic abnormalities on as many of these as possible and have analysed an initial cohort of 1565 workers aged over 70 years. 1383 (88%) of these subjects started work underground before the age of 20, and their median duration of underground work was 39 years (1–50); 1377 (90%) reported cough and sputum and 1526 (99%) reported breathlessness, 812 (52%) used inhalers, 1145 (75%) were current or ex-smokers with a median reported cigarette consumption of 10/day, 514 subjects (34%) had FEV1 values of more than 1 litre below the predicted value. Of these, 147 (29%) had radiographic evidence of pneumoconiosis and received compensation. The median disability assessment was 50% (10–100%). Stepwise linear regression analysis suggested that FEV1 was positively related to height and was negatively related to age, the use of inhalers, smoking, and the presence of cough. There was a slight but significant (p<0.001) positive relationship with years of underground work with FEV1, increasing by 0.0071 per year of underground exposure.

Work-related symptoms and specific sensitisation in two occupational cohorts

P CULLINAN, D LOWSON, M NIEUWENHUIJSEN, S GORDON, C SANDFORD, RD TEE, KM VENABLES, JC MCDONALD, AJ NEWMAN TAYLOR Department of Occupational and Environmental Medicine, National Heart and Lung Institute, London The development of symptoms and specific sensitisation over up to five years has been compared in two newly exposed occupational cohorts; incident, work-related respiratory, eye/nose or skin symptoms and positive, specific skin prick tests (SPT) were related to measured whole shift exposures to aeroallergen or dust in 208 laboratory animal workers and 103 flour exposed workers. In both cohorts the incidence of all symptom types was highest in the first year of exposure; eye/nose symptoms were most commonly reported. Amongst flour workers there was no evidence of a relationship between new symptoms and a positive SPT to flour; in laboratory workers with specific sensitisation and symptoms these tended to be identified simultaneously. In the flour workers an exposure response relationship was seen only with exposure expressed in terms of dust; there were no associations with atopy or smoking. In laboratory employees regression analysis revealed an association between eight hour rat urinary aeroallergen exposure and new skin (OR=1.12 per 10 μg/m3) and eye/nose symptoms (OR=1.01), and the development of a specific SPT (OR=1.13); smoking and asthmatoid status were also associated with these outcomes. These findings are consistent with those from an initial cross sectional analysis and suggest that the development of work-related disease in these two groups may reflect different pathological processes or selection pressures.

Comparison of across shift and across week variations in lung function in symptomatic textile workers and matched controls

GJ WARBURTON, AM FLETCHER, CAC PICKERING, RM NIVEN, LA OLIDHAM, JC FRANCIS Northwest Lung Centre, Wythenshawe Hospital, Manchester Eighty textile workers with one or more respiratory symptoms and 84 asymptomatic control workers (defined by questionnaire) performed spirometric and bronchial reactivity (BR) testing at the beginning and the end of the first and fourth shifts of the working week. The symptomatic workers were significantly older than the controls, and demonstrated lower levels of percent predicted lung function (p<0.05 only for FEF25–75). There was no difference between the groups for current or past cotton dust exposure. On day 1 both groups demonstrated falls in spirometric values and increases in BR across the shift. None of the differences were significant. On day 4 both demonstrated a fall in FEV1, and FCC with rises in FEF25–75 and FEF25–65. There was a significant increase in BR in the controls as opposed to the symptomatics on this day (p<0.05). The symptomatic workers demonstrated significantly greater falls in FEF25–65 across the working week than the controls. There was little difference for FEF25–75 and FEF25–65. Exposure to textile dust seems to have similar effects upon workers with and without symptoms, and this effect is more marked on the first working day. Symptomatic operatives
do not recover lung function as quickly as the controls leading to declines across the working week. Asymptomatic workers seem to experience the largest changes in BR.

Are the respiratory health effects seen in ceramic fibre manufacturers due to the dust rather than the fibre exposure?

P S BURGE, WN TREATHOWEN, JM HARRINGTON, I A CALVERT  Birmingham Heartlands Hospital and Institute of Occupational Health, Birmingham University, Birmingham We have carried out a cross sectional survey of all seven European ceramic fibre manufacturing plants, estimating current exposure using full shift personal samplers, measuring insensible dust and respirable fibre levels. The response rate was 89% of 708 current employees. Plant and job specific cumulative exposure levels were calculated from detailed occupational histories based on cumulative exposure. Respiratory health was assessed by questionnaire and spirometry. Odds ratios relating current symptoms to current exposures were calculated by multiple logistic regression, and regression coefficients for lung function related to cumulative exposure were calculated using multiple linear regression, each controlling for the effects of respirable fibre and insensible mass exposures separately and together, after adjusting for age, sex, plant (and hence country) and smoking. Significantly increased odds ratios for current exposure to both dust and fibres were seen for dry cough, stuffy nose, eye and skin irritation and breathlessness. There were significant decrements in FEV\(_1\), FEF\(_{25-75}\) and a few other fibres seen in current smokers which were reduced to non-smokers and not in fibres exposed to dust. No effect was seen in lifelong non-smokers, intermediate effects were seen in ex-smokers. In conclusion, current symptoms were related to both insensible mass and respirable fibre exposures. Dose-related decrements in lung function related only to cumulative respirable fibre exposure and were confined to smokers, suggesting a promoting effect of ceramic fibres on the effects of cigarette-related declines in FEV\(_1\).

Scottish national survey of tuberculosis notifications in 1993 with special reference to the prevalence of HIV seropositivity

AG LEITCH, M RUBILAR, GI FORBES, J CURNOW, G BOYD, S BURNS, B WATT  Tuberculosis Survey Study Office, Royal Victoria Chest Clinic, Chalmers Hospital, RIE NHS Trust, Edinburgh All notifications of tuberculosis in Scotland in 1993 were surveyed by questionnaire and a 100% response rate was achieved by supervising clinicians. HIV testing was requested for under 65 year olds. Of 497 notifications, 423 (85%) were white and 58 (12%) were from the Indian subcontinent (ISC). Of those from the ISC 85% were aged under 55 years whereas 64% of white patients were aged over 55 years. 37% of white and 47% of ISC patients were male. Pulmonary disease was found in 74%, non-pulmonary in 22%, and combined disease in 4% of patients. ISC (47%) had more non-pulmonary disease than white (15%) patients. Genitourinary disease (33%) and pleural effusion (17%) were increased in ISC patients. Cancer and histological disease was (67%) in ISC patients. Cultures for tuberculosis were positive in 64% of white and 65% of ISC pulmonary cases with 48% and 45% positive smears. Of 244 HIV tests performed three, including one previously known, proved positive. Five other HIV positive patients were known, giving an HIV positivity rate of 1-6% for the 497 Scottish tuberculosis notifications in 1993. Four HIV positive patients were African. The annual tuberculosis notification rate for Scotland in 1993 was 9.7-10\(^3\). In white patients the rate was 8.4-10\(^3\) with higher rates for men and increasing rates with age in both sexes. The ISC population rate for Scotland was 179.8-10\(^3\). Notification rates for white patients were higher in Glasgow (13.0-10\(^3\)) than in Edinburgh (6.9-10\(^3\)). Tuberculosis notifications in Scotland have fallen by 5% since 1992. HIV infection is an uncommon accompaniment of tuberculosis disease in Scotland.

Changing pattern of respiratory tuberculosis in adult patients from the Indian subcontinent

OR McCARTHY, LP ORMEROD  Newham Chest Clinic, London; Blackburn Royal Infirmary, Blackburn, Lancashire Clinical observation over 10 years in both clinics in high tuberculosis prevalence areas suggested an altering pattern of respiratory tuberculosis in adult patients from the Indian subcontinent (ISC), with a declining proportion of isolated mediastinal lymphadenopathy, and an increasing proportion of smear positive and culture positive disease. Details of all 878 adult ISC (age >15) cases treated in the two centres in the years 1981-92 were analysed. The main site of disease, spumtum status at diagnosis, time since first entry to the UK, ethnic group, and sex were known for all patients. General logistic regression showed a highly significant effect by year (p=0.0002). There was a highly significant trend over time (p=0.0002), with an increasing percentage of cases with increased percentage culture positive disease (p=0.47). There was also a significant trend to decreasing percentage mediastinal lymphadenopathy over time (p=0.002), with no difference between centres (p=0.65) or centre by year interaction (p=0.47).

Results of the national survey of tuberculosis notifications in England and Wales in 1993

J M WATSON ON BEHALF OF A PHLS/BTS/DH COLLABORATIVE GROUP  PHLS Communicable Disease Surveillance Centre, London A survey of tuberculosis notifications in England and Wales in 1993 was carried out to determine the occurrence of the disease in the whole population and to distinguish the disease in known demographic subgroups, from the first six months of the survey on site of disease and bacteriological results. The prevalence of HIV infection in adults (16-54 years) notified over the whole year was estimated using unlinked anonymous HIV testing. By 30 September 1994, for 2338 eligible notifications in the first six months, 1997 (85%) clinical forms had been received. Of these, 41% patients were white, 43% from the Indian subcontinent (ISC), and 16% from other ethnic groups (the equivalent figures in the 1988 survey were 53%, 39%, and 8%, respectively). After exclusion of previously treated cases and allowing for incomplete data, the crude annual notification rate for England was 8.7 per 100 000 compared with 8.6 in 1988. The rate in the white and ISC ethnic groups was 3.9 and 11.6 per 100 000 compared with 4.7 and 120 in 1988, respectively. Bacteriological results were available for 1029 (88%) of the 1169 pulmonary cases: 659 (64%) were positive on culture; 574 (56%) had positive smears. Of the 659 with positive cultures (41%) were resistant to one or more first line drugs, including 12 (1.8%) resistant to isoniazid alone and a further 13 (2.0%) to isoniazid in combination with other drugs. Specimens were received for 1073 (40%) of the 2701 patients eligible for inclusion in the HIV prevalence survey: 22 (2-1%) were positive. The continued decrease in white and ISC rate of infection is felt to be due to improved diagnostic services and education, although the demographic structure of the population and the increased number of cases in the other ethnic groups have contributed to the overall increase in notifications.

UK mycobacterium resistance network

D BENNETT, J WATSON, D KUMAR, M YATES, T JENKINS  PHLS Communicable Disease Surveillance Centre (CDS), Dulwich Public Health Laboratory, Mycobacterium Reference Unit In order to characterise drug resistance trends for Mycobacterium tuberculosis isolates from the UK, which are currently frequently than under the five yearly national surveys, the PHLS Mycobacterium Reference Unit and Regional Tuberculosis Centres have agreed to report information on mycobacterial isolates they receive for UK residents to CDS. The reference laboratories in Scotland and Northern Ireland, as well as the Wombourough Hospital Laboratory, are also participating. An electronic system for regular reporting of the relevant demographic, geographical, specimen site, sputum microscopy, and antibiotic susceptibility information, based on an agreed minimum data set, has been piloted since January 1994. Resistance results are made on all drug sensitive isolates and isolates in which susceptibility has changed. A preliminary analysis of 643 deduplicated reports received on M tuberculosis isolates through April 1994 indicates that resistance to isoniazid and rifampicin combined could be rising in England and Wales: combined resistance was 1-8% compared with a preliminary estimate of 0.8% from the 1993 National Survey of Tuberculosis Notifications in England and Wales, and 0-6% from the 1988 survey. Increases were also seen in other resistance measures. Resistance patterns by disease site did not differ significantly among ethnic groups, except that significantly more pulmonary isolates as...
were notified, positive of Proceedings (59%) whether the sets the planning the London Hospital, London; be HIV negative, negative, or of Medical of a BAL longer and date records. In 1992-3. Ten cases were notifiable and diagnosis was established by either a 15% diurnal variability in FEV₁ or a >15% bronchodilatation to inhaled salbutamol. Patients were randomised to one of the following drug classes: a short acting β₂ agonist (Aerol in Autohaler 200 µg pm or Bricanyl turbohaler 0·5 mg pm); a long acting β₂ agonist (Serevent 50 µg bd); an inhaled steroid (Becotide 200 µg bd or Pulmicort Turbohaler 100 µg bd); nedocromil sodium 4 mg qds; and oral theophylline (Theo-maphine 250 mg bd or Theodur 300 mg bd). Theophylline levels were monitored to achieve levels of 10–20 mg/l. Symptoms and daily FEVR measurements were recorded in a diary card. Baseline characteristics of the five groups were similar: mean FEV₁ (predicted) 2-61 (82%), FVC 3-61 (91%). Wheeze, cough and expectation were present on 4-2, 3-8, and 2-8 days per week. At other month’s follow up improvement in number of symptom-free days was seen in the group given inhaled steroids. Mean days per week with wheeze fell by 1-3, cough by 0·5, and expectation by 1·5. Nedocromil sodium produced similar but less striking results of 0-8, 0-3, and 0-8 respectively. Other modalities of treatment produced no significant change in symptoms. In this group of mild asthmatics mean improvements in FEVR, was greatest in the steroid group (11%) followed by the nedocromil (9%) and salmeterol (7%) groups. There was no change with short acting β₂ agonist, or theophylline. Anti-inflammatory therapy produces the greatest symptomatic and physiological improvement in mild asthma. [Supported by Astra, Fisons, Allen & Hanburys, and 3M.]

Patterns of prescribing of inhaled steroids over a seven year period in a general practice and its implications

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There is evidence that the prevalence of asthma appears to be increasing (Bur et al. Arch Dis Child 1989;64:1452–6). As asthma increases so does the burden of asthma costs. It is recommended that an essential part of the management of asthma is the use of inhaled steroids. An audit was carried out in my practice to look at the increase in prescribing costs and its relation to both the drugs used and the costs per patient. Between 1987 and 1994 the recorded asthma population in my practice of 10 500 patients has increased from 345 to 1106 (10.5%). A variety of factors may be responsible: increasing diagnosis and prescribing costs, prescribing patterns (0.85) to changing local, population, and asthmatics changing practices (30% of transferring patients). Asthma prescribing costs have more than doubled over the study period from £39 706 in 1987–8 to £89 040 in 1993–4. This consists mainly of spending on inhaled steroids which has risen from £4 560 to £25 640 with bronchodilator costs and falls for other asthma medications. The number of asthmatics using inhaled steroids has increased from 110 to 694. Thus, annual drug costs per diagnosed asthmatic have fallen from £115 to £80 with inhaled steroids staying between £40 and £48. Clinical improvements have been found in terms of reduced total admissions for asthma over the age of three (43%) and outpatient attendances (65%) over the period. For individual asthmatics changing to inhaled steroids the decrease is 74% and 90% respectively. Total recorded exacerbations fell by 25% in the last two years. The data in this study support the premise that increased use of inhaled steroids in the management of asthma can be cost effective.

Tuberculosis: how to improve notification rates

JS BROWN, F WELLS, G DUCKWORTH, NC BARNES The London Chest Hospital, London; The Royal London Hospital, Whitechapel, London Tuberculosis (TB) is a statutory notifiable disease. The accuracy of the statistics from notifications is important for the planning of TB services and for thorough contact tracing. However, we have previously shown over a quarter of cases of TB were not notified for the years 1985–9 in an east of London served by the London Chest and Royal London hospitals (Sheldon et al. Thorax 1992;47:1015–8). As a result several changes in practice were instigated to improve notification rates: clinicians were encouraged to refer all cases of TB to the chest physicians for advice on management; suggestive history reports and all positive microbiology reports were sent to the communicable disease control (CDC) consultant to check whether the patient had been notified or not; and pharmacy informed the CDC consultant about patients prescribedisoniazid. We now report a repeat audit of TB notification rates for the years 1992–3 from these two centres. Identical methods were used to identify cases as the original study, and comparisons were made between the two sets of data. 252 cases of adult TB were identified for the years 1992–3, giving an annual incidence of 126 cases/year which is similar to the original study (122/year). The mean age was 46 years and 148 (59%) cases were male. There was a significant increase in the proportion of patients of African origin from 15 out of 580 (3%) in the original study to 29 out of 252 (12%, χ² 29-5, p<0.0001) for cases diagnosed in 1992–3. Ten patients (4%) were known to be HIV positive, whereas only four patients (0-7%) were known to be HIV positive in the 1984–9 group. 236 cases (94%) were notified, a highly significant increase of 21% over the original study where 73% of cases had been notified (χ² 42-9, p<0.0001, confidence intervals 16% to 25%). Only one smear positive patient was not notified. We have therefore shown that changes in the practice of notification for TB can lead to a substantial improvement in notification rates and so provide accurate data for contact tracing and on the local incidence of TB.
have studied serum osteocalcin and serum parathyroid hormone (PTH) pre and post oral calcium load (1200 mg calcium) in 10 male asthmatic patients (mean age 52-7 years) taking inhaled BDP (mean dose 1000 mg per day) and 10 asthmatic male patients (mean age 55-6 years) on inhaled BDP with oral corticosteroid mean daily dose prednisolone 9 75 mg (P). These groups were compared against 10 aged matched male controls (mean age 54-4 years) on no corticosteroid preparation (C). Subjects on steroid preparations had been taking their medication for at least three months with no change (P 0-25-10 years, BDP 0-25-7 years). All subjects had normal renal and bone biochemistry. Osteocalcin levels were all in the normal range and there was no difference statistically between groups (C 5-50-27; P 4-31 (1-57); BDP 5-74 (3-22)). PTH levels were all in the normal range pre calcium load (C 4-54 (1-30) pmol/ml; P 4-16 (0-86); BDP 4-04 (1-28)). PTH decrement after calcium load (C 2-02 (1-35) pmol/ml; P 1-81 (0-75); BDP 1-64 (0-76)) were all in the normal range suggesting calcium clearance. Both patients and controls on chronic stable corticosteroid use and controls did not have different osteocalcin levels, PTH levels and response to calcium absorption. These findings indicate stable bone biochemistry in patients with chronic stable steroid ingestion and suggest interpretation of findings of acute studies should be made with caution.

**Inhaled corticosteroids, asthma, and long term bone metabolism: collagen crosslinks, osteocalcin, and densitometry after one year**

K ANDERSON, SJ GALLACHER, WD FRASER, T SPEEKENBRINK, D VERNON, IT BOYLE Department of Respiratory Medicine, Western Infirmary; University Department of Medicine, Royal Infirmary, Glasgow Hitherto, most studies of bone metabolism in asthma relating to inhaled corticosteroids have been cross sectional, of short duration, and may have included patients who have taken oral corticosteroids. We compared the effect of inhaled beclomethasone (BDP) 500 mg bd (n = 5, mean age 50 years) and inhaled budesonide (BUD) 400 mg bd (n = 4, mean age 41 years) on bone metabolism (urinary collagen crosslink excretion, osteocalcin, and lumbar spine/hip density by dual energy x ray absorptiometry) measured pretreatment and after one year in patients with asthma who had never taken oral corticosteroids. The drugs were inhaled through a spacer followed by rinsing the mouth with water in an attempt to reduce non respiratory drug deposition. Mean (SE) pretreatment L2-4 densities were similar in both groups (BDP 1.162 (0.020) g/cm² v BUD 1.179 (0.030), p = NS). However, L2-4 density fell significantly over the year after BDP (5-8 (3-1) % v BUD +1.9 (1-5)%), p = 0-028. No change was noted in femoral neck density. Pretreatment serum osteocalcin was similar (BDP 8-0 (2-0) ng/ml v BUD 8-0 (2-0) BUD, p = NS) and did not change significantly after one year. Pretreatment serum osteocalcin was similar both to pyridinium/creatinine (BDP 12-8 (1-0) v 15-2 (1-9) BUD, p = NS) and free deoxypyridinium/creatinine (BDP 2-5 (0-2) v BUD 2-9 (0-3), p = NS) were not changed with treatment. These results suggest that, although biochemical markers of bone turnover may show no change following treatment with BDP or BUD, lumbar spinal density would appear to fall after one year of BDP. Urinary collagen crosslink excretion is the most sensitive biochemical marker of bone resorption, however, given our results and the relative precision of crosslink measurement of 11% (Gallacher et al. Eur J Endocrinol 1994) and 0-8% for bone densitometry, the latter may be more useful in clinical practice over the long term.

**Markers of bone metabolism in asthmatic children receiving inhaled corticosteroids**

JOLO JM DOULL, M WHITE, NICHOLAS J FREEZER, STEPHEN T HOLGATE University Medicine, Centre Block, Southampton General Hospital, Southampton Osteocalcin is produced almost exclusively by osteoblasts, and as such is a sensitive index of bone formation. We quantified inhaled corticosteroids by hydroxyproline, which is present in urine, and is thus a measure of bone resorption. To determine the effect of inhaled corticosteroids on bone turnover in children we measured serum osteocalcin and urinary hydroxyproline in a group of 94 mildly asthmatic children receiving inhaled beclomethasone dipropionate (BDP) 200 mg twice daily as a dry powder in a randomised, double blind, placebo controlled manner. Samples were collected prior to starting BDP and after three and six months treatment. Serum osteocalcin was measured by radioimmunoassay. An overnight urine was collected and urinary hydroxyproline measured as per Podenphant et al, and expressed as a hydroxyproline/creatinine ratio (Hyd/Cre). There was no significant difference in osteocalcin at any of the three timepoints between the BDP and placebo treated children (baseline: 19.3 ± 21.44 µmol/l, p = 0.12; three months: 18.34 ± 20.20 µmol/l, p = 0.12; six months: 15.51 ± 17.69 µmol/l, p = 0.19), or in Hyd/Cre at any of the three timepoints (baseline: 17.8 ± 19.5 µmol/ mmol, p = 0.45; three months: 13.2 ± 18.2 µmol/mmol, p = 0.22). Serum osteocalcin was significantly correlated at three (r = 0.5, p = 0.001) and six months (r = 0.53, p = 0.001) with growth over the six month period. At conventional doses BDP does not affect serum osteocalcin or urinary Cre/HD in mildly asthmatic children.

**Suppressive effects of therapeutic doses of fluticasone proponate on the HPA axis in healthy volunteers**

A GRAHNÉN, A LING-ANDERSSON, RM BRUNDIN, SK ECKERNÄS Pharmacoe Medical Consultants, PM AB, Upsala, Sweden A new inhaled glucocorticosteroid, fluticasone proponate (FP), has recently become available. It has been claimed that FP has no clinically significant effects on the hypothalamo-pituitary-adrenal (HPA) axis, primarily based on data from single, morning plasma cortisol assessment. In two studies we have investigated the effect on the HPA axis following single and repeated doses of inhaled FP using a multiple sampling technique. Study 1: FP was administered as single doses (250, 500, and 1000 µg) respectively) and repeated doses (1000 µg bid) from Diskhaler. A single dose resulted in a small suppression of cortisol, but did not affect cortisol suppression after calcium load. After repeated dosing, no cortisol suppression was significantly different from placebo. After seven repeated doses of fluticasone (100 µg bid), the cortisol suppression was 65%. Study 2: FP from Diskhaler and BUD from Turbuhaler were given as single doses (1000 µg and 800 µg, respectively) and repeated doses (1000 µg bid and 800 µg bid, respectively). The study was open, randomised, placebo controlled, crossover design. Twenty five healthy male volunteers took part in the study. For single doses of FP, the cortisol suppression, measured as AUCs, was 8%, 19% and 29% for 250 µg, 500 µg and 1000 µg respectively. The single dose resulted in a small suppression of 16%. Cortisol suppression, after all single doses, was significantly different from placebo. After seven repeated doses of fluticasone (100 µg bid), the cortisol suppression was 65%. Study 2: FP from Diskhaler and BUD from Turbuhaler were given as single doses (1000 µg and 800 µg, respectively) and repeated doses (1000 µg bid and 800 µg bid, respectively). The study was open, randomised, placebo controlled and crossover. Twenty four healthy male volunteers participated. In both studies the single doses were administered at 10.00 pm and the repeated doses at 10.00 pm and 10.00 am for 3-5 days. Multiple plasma cortisol samples were taken during a 20 hour period after each single dose and after the last (7th) repeated dose. Single doses of FP (1000 µg) and BUD (800 µg) resulted in a cortisol suppression of 25% and 26%, respectively. Repeated dosing for 3-5 days resulted in a cortisol suppression of 55% for FP (1000 µg bid from Diskhaler) and 54% for 800 µg bid from Turbuhaler. The difference was statistically significant (p<0.001). We conclude that FP from Diskhaler, within the therapeutic dose range, results in a significant suppression of plasma cortisol which is more marked after repeated dosing.

**Genioglossus muscle response to negative airways pressure in awake humans: lack of effect of stimulus timing within the respiratory cycle**

P NOLAN, DM O'LEARY, SA O'NEILL, SM ROE, BG O'REGAN, WT MCGHAN Department of Respiratory Medicine and Physiology and the Respiratory Sleep Laboratory, University College and St. Vincent's Hospital, Dublin, Ireland Upper airways (UA) negative pressure elicits a reflex increase in UA dilator muscle activity to help maintain a patent airway. Studies in anaesthetised rabbits have shown that the genioglossus (GG) response to UA negative pressure is greatest during early inspiration, protecting the UA at the time in the respiratory cycle when it is most vulnerable to collapse (Woodall et al. J Appl Physiol 1989;67:366-70). The purpose of this study was to determine if the UA negative pressure responses to UA negative pressure is influenced by the timing of the pressure stimulus within the respiratory cycle. We studied six healthy naive volunteers with their informed consent. The subjects lay supine and breathed through a nose mask. GG EMG was recorded using bipolar intramuscular surface electrodes. Pulses of negative pressure (350 ms – 30 cm H2O or 30 cm H2O) were applied to the nose mask at eight times in the respiratory cycle, ±100 ms, ±300 ms, ±600 ms, and ±1000 ms after the onset of inspiration (INS) and ±100 ms, ±300 ms, ±600 ms, and ±1000 ms after the onset of expiration (EXP). The GG EMG
reflex activity in comparison with the antagonist muscle LP, using electromyography in seven patients with sleep apnoea. Results were expressed as % maximum EMG activity in response to upper airway negative pressure application via nose or mouth. PP demonstrated phasic inspiratory activity during nose and mouth breathing. Application of negative pressure via the mouth caused a significant increase in IP activity (p<0.02) reaching 70-90 minutes, at -12.5 cm H2O compared with 87% for LP (p<0.001). Negative pressure applied via the nose caused only a non-significant trend to increased activity (p=0.13) reaching 60% maximum at -12.5 cm H2O for PP compared with 80% for LP (p<0.001). This differential response to nasal negative pressure application for LP and PP may promote retropharyngeal obstruction in patients with sleep apnoea.

Effect of sleep fragmentation on daytime function

SE MARTIN, TJ DABRY, NJ DOUGLAS Respiratory Medicine Unit, Department of Medicine and Department of Psychology, University of Edinburgh, Edinburgh Patients suffering from the sleep apnoea/hypopnoea syndrome (SAHS) show impaired daytime function and decreased sleep latencies on the MSLT. To help understand whether these features of SAHS are caused by sleep fragmentation or hypoxia, we have studied the effects of sleep fragmentation alone on daytime function in normal subjects. Nine normal subjects were studied over four nights and two days in the sleep laboratory. The nights were divided into two pairs. The first of each pair was for acclimatisation, and on the second the subject either slept undisturbed or had sleep fragmented with SBP pulses every 90 minutes. Sound volume was titrated to cause a return to theta or alpha electrocortical activity for a minimum of three seconds but no longer than 15 seconds. Both the second nights were followed by a full day's testing of daytime function using psychometric tests, the MSLT and the Maintenance of Wakefulness Test (MWT). Total sleep time did not differ between study nights (mean (SD) 400 (20) min undisturbed; 392 (19) min disturbed, p=0.5). However, sleep architecture was altered during the fragmented night with significant increases in stages 1 (p=0.02) and 2 (p=0.002) and decreases in stage 4 (p=0.001) and REM (p=0.03). The time to REM onset was delayed on the fragmented night (p=0.04). Mean (SE) sleep latency decreased on MSLT (10 (1) 7 (1) min, p<0.03) and MWT fell (29 (3) 20 (3) min, p<0.01) after fragmentation. Subjects showed a decrease in their levels of energetic arousal (p=0.01) and an increased tense arousal (p<0.05) on the UWIST scale after sleep fragmentation. One night of sleep fragmentation adversely affects the ability of normal subjects both to stay awake and to fall asleep. Energetic arousal is decreased and tense arousal is increased after a fragmented nights sleep. Their ability to perform other tasks of mental flexibility are not altered by sleep fragmentation in this ongoing study.

Evaluation of pulse transit time as a screening respiratory sleep study

DJ PITSON, JR STRADLING Oster Chest Unit, Churchill Hospital, Oxford Two important components of a respiratory sleep study are a measure of inspiratory effort (marker of upper airway narrowing) and a measure of arousals (marker of sleep disturbance). In an all night beat-to-beat blood pressure (BP) profile, the size of the falls in systolic blood pressure (SBP) with inspiration (pulsus paradoxus) correlates well with the degree of inspiratory effort and there are rises in SBP with each arousal. Pulse transit time (PTT), measured from the ECG R wave to detection of the pulse shock wave at the finger, follows changes in BP well. The size of the inspiratory rises in PTT correlate well with inspiratory effort and falls in PTT are sensitive markers of arousal. Unlike SBP, PTT can be measured noninvasively using a portable device. Twenty six patients presenting to the sleep unit for investigation of a possible sleep-related breathing disorder had a domiciliary PTT study and a laboratory PTT recording in conjunction with a polysomnography sleep study. PTT records were analysed to calculate the mean size of the inspiratory rises in PTT for the night and the number of arousal-related falls in PTT/hour (r=0.91). In the eight normal patients (no obstructive sleep apnoea (OSA) or snoring) the average inspiratory PTT rises ranged from 7 to 14 ms and in the nine patients subsequently established on nasal continuous positive airway pressure (NCPAP) for OSA PTT rises ranged from 14 to 39 ms.
PTT arousals ranged from 2 to 15/hour in the normal subjects and from 15 to 94/hour in the NCPAP group. Thus, PTT data appear reproducible and capable of differentiating between two groups of patients defined as normal or requiring NCPAP on clinical history and polysomnography. PTT may therefore be a useful tool in a screening respiratory sleep study.

Survey of pneumothorax management in Wessex

E Neville, J Turner, D Lipscombe, R Lea, J Waddell and A Roberts On behalf of the Wessex Thoracic Physicians and Surgeons Chest Clinic, St Mary’s Hospital, Portsmouth A survey of current clinical practice was conducted on 169 patients admitted in Wessex between 1 January 1992 and 1 July 1992. Forty four (26%) were managed by initial observation from one to 66 days, and only six patients had either aspiration or tube drainage at a later stage. Eighteen (11%) were treated by initial aspiration. In nine of these a subsequent intercostal tube was inserted. One hundred and seven (63%) patients were managed by initial intercostal tube drainage. Instructions about not clamping the tube were written in the notes in only 48 (40%) of patients. Thirty four (30%) patients had suction applied and, of those, only two were recorded as having a satisfactory apparatus applied.

After the tube was removed, there was incomplete expansion of the lung in 28 (29%) of cases. There were nine deaths, six with underlying respiratory disease, two of whom were under the age of 65. Surgery was performed in 22 (13%). A 20 per cent return to work and school (for example, not flying) was recorded in only 36 (22%) of patients.

We conclude that (1) the BTS guidelines do not adequately deal with removal of chest drains; (2) the management of chest drains and suction is unsatisfactory; and (3) advice to the patient at discharge was inadequate. We recommend that (1) each junior doctor’s induction package should include four algorithms or lists: (a) management decisions, (b) management of chest drains, (c) when to refer for specialist respiratory advice, (d) discharge checklist; and (2) each ward where a chest drain may be cared for should have a wall suction adaptor with clear instructions as to its use.

Audit of inpatient bronchoscopy requests: should hospital physicians have open access?

PH Johnson, H Hopkinson, IDA Johnston, WM Kinneir Queen’s Medical Centre, University Hospital, Nottingham Open access to upper gastrointestinal endoscopy is commonplace, but it is traditional in most hospitals for a chest physician to assess referrals for bronchoscopy beforehand. This latter convention is time consuming. We have audited requests for bronchoscopy on inpatients to see if an open access service would be cost effective. Of 102 patients referred for a respiratory opinion over a period of six months, 18 involved a request for bronchoscopy. Nine were from medical specialities, seven from geriatrics, and two from surgical wards. All referrals were seen by a registrar or consultant in respiratory medicine. Of the 18 requests, 10 were considered to be appropriate by the respiratory physician who saw the patient, but four of these 10 were unfit for the procedure. In the other eight patients the referring clinicians had unrealistic expectations of the ability of bronchoscopy to yield the information that they required. Had an open access policy been in place in our hospital, 12 of the 18 patients referred would have had the procedure cancelled on arrival in the bronchoscopy room, entailing wasted bronchoscopy time and money, and distress to the patients. Even in those cases in whom bronchoscopy was subsequently performed, assessment of the patients beforehand by a respiratory physician was often valuable, for example in making provisional management plans.

We conclude that all patients being considered for bronchoscopy should be seen by a respiratory physician, and that the time involved in these assessments is cost effective. Open access to bronchoscopy for hospital physicians might be appropriate if they were issued with guidelines on the indications and required fitness.

Cigarette smoking and death certification

TJ Fletcher, RC Horton, JG Ayres Chest Research Institute, Birmingham Heartlands Hospital, Birmingham; Cardiac Transplant Unit, Queen Elizabeth Hospital, Edgbaston Since 1992 doctors have been able to record cigarette smoking as a cause of death on death certificates without reporting the death to the coroner. In the 10 years prior to 1992 only some 200 such cases had been reported to the coroner countrywide despite an estimated 110 000 deaths per annum caused by smoking related disease (BMJ 1992;305:552). This new measure was taken in part to improve the accuracy of statistics on smoking-related deaths. We have gained the impression that, despite the new regulation, smoking is rarely recorded on death certificates. A retrospective analysis was undertaken of all deaths occurring in Birmingham Heartlands Hospital in January 1994. Age, sex and cause of death as recorded on the death certificate counterfoil was recorded in each case. One hundred and thirty two deaths were reported during this period (74 male, 78 female, mean age 75, range 19–105). None had smoking recorded as a cause of death. Hospital notes were sought and details of smoking habit recorded. To date we have obtained 72 case notes for analysis. Of these, 32 (44%) had inadequate documentation of smoking history in that we were unable to determine from the notes whether the patient had ever been a smoker. Five more (7%) were comatose on arrival and unable to give smoking details, and nine (13%) were non-smokers or had not smoked for at least five years. The remaining 26 (36%) were still smokers at the time of hospital admission or had smoked within the last five years. From the causes of death recorded in this group 21 were considered to have died from a condition recognised to be smoking related (nine from bronchial carcinoma, five from chronic obstructive pulmonary disease or emphysema, four from ischaemic heart disease, two from oesophageal carcinoma, and one from a stroke). We conclude that smoking is still not recorded on death certificates despite regulations designed to improve the accuracy of information regarding smoking-related deaths.

Variability in the hospital management of acute emergencies and implications for future savings

H Moodgil, J Hammersley, M Anderson, AG Leitch Respiratory Medical Unit, City Hospital, RIE NHS Trust It is assumed that the cost of care for a given patient is determined by the severity, complexity, or duration of the patient’s illness with little attention paid to physician behaviour in this setting and despite variability in clinical practices.

Four hundred and thirteen acute medical admissions to a specialist respiratory medical unit were audited to document the overall and diagnosis-related (DR) patterns of investigation (laboratory and radiological), duration of admission, and patient outcome (disposal) for the five consultant teams: team A (99 patients), team B (98), team C (98), team D (97), and team E (89).

The variability of investigation (laboratory and radiological) was determined for team A. It was noted that approximately 40% of patients had one or more investigations (laboratory and radiological) ordered on admission, and that 74% of patients were investigated by at least two different consultants. The team A investigation policy was critically examined, and the variability was reduced to a minimum. There was no increase in the rate of admissions, hospital stay, or mortality. The audit concluded that the variability was the non-essential, and represented areas in which savings could be made. Objectives for testing differ and although it is not possible or even desirable to eliminate all unnecessary investigations the cost implications cannot be ignored. Investigations should be directed to where the result, even if normal, is important in the management of the patient. A change in physician behaviour is proposed which should include a regular assessment of “why” and “to what purpose” investigations are performed.
Audit of discharge rates in respiratory outpatient clinics: NHS versus the private sector

JW WALES, E DRAPER The Glenfield Hospital NHS Trust and Department of Epidemiology and Public Health, University of Leicester, Groby Road, Leicester Outpatient referrals consume an increasing proportion of medical manpower. We have set up a study to evaluate the difference in follow up rates between private (consultant only) clinics and NHS clinics held during the same period. All new patients seen in one consultant's private practice in 18 months (n = 372) and two simultaneous NHS clinics (n = 527) were entered into the study. Demographic data, conclusions and reasons for non-attendance were recorded. Findings by coded ICD-9, were recorded. Follow up continued for a further 18 months. All private and all but five NHS patients were seen by one observer at the first consultation. Follow up was continued by the same observer in private practice and usually by registrars and senior house officers in the NHS sector. A marked difference in social group existed (social group I and II: NHS n = 109, 20.7%; private sector n = 178, 47.8%, p<0.0001). The table shows numbers of patients attending at six monthly intervals after entry into the study was closed: 317 private (85.4%) and 340 NHS (64.5%) were discharged by the second follow up appointment. There was an excess in the NHS sector of patients with chronic airways limitation and lung cancer. Authors were bound to present their conclusions in this form, for which reasons for such a marked variation in follow up rates are discussed. We feel that the study supports the value of consultant only clinics in the NHS to support patients who may be more socially disadvantaged or sicker than those referred privately.

Characteristics of infants ventilated with fast rates who receive muscle relaxants

DP COCHRAN, NJ SHAW Neonatal Unit, Liverpool Maternity Hospital, Liverpool With the increasing use of fast ventilator rates (60-80 breaths/min) fewer preterm infants appear to require muscle relaxants. However, even with faster rates there remain a group of babies for whom they are prescribed on subjective, clinical criteria. We studied 97 infants (birth weight <2kg), ventilated using rapid rates, of whom 24 (24.7%) received muscle relaxants as judged necessary by the attending clinicians. There was no significant difference in birth weight or gestation between the two groups. Infants who were paralysed had markedly more severe lung disease with higher inspired oxygen concentrations (median 1.0 v 0.65, p<0.0001) and peak inspiratory pressures (median 27.5 v 22.0, p<0.0001) on day 1. The paralysed group suffered high mortality (10/24 v 8/73, p<0.001), a higher incidence of parenchymal cerebral haemorrhage (6/25 v 4/72, p<0.05), required inotrope more frequently, (13/23 v 19/73, p<0.05) and had a higher incidence of pneumothorax (6/25 v 10/72, NS). The risk of death or chronic lung disease was higher in the paralysed group but was only significant at 36 weeks gestation (17/25 v 28/72, p<0.05) and not at 28 days of age. These data suggest that infants who receive muscle relaxants are likely to have severe lung disease and ultimately a poor prognosis. There are a number of possible explanations for these associations. We believe these data should prompt a re-examination of the use of muscle relaxants in the era of fast rate ventilation to ensure that their prescription is appropriate and that they are not contributing to the poor outcome in this group of babies. Newer strategies of ventilation may hold more promise for these patients.

Value of flexible fiberoptic bronchoscopy in children with persisting lobar collapse

DE LACY, RL SMYTH, DP HEAFF Respiratory Unit, Royal Liverpool Children's NHS Trust, Alder Hey, Liverpool Flexible fiberoptic bronchoscopy has been available at Alder Hey Hospital since 1992. We have reviewed the results of bronchoscopy in children who presented with persisting lobar collapse (despite conventional treatment of two weeks antibiotics and physiotherapy) from January 1992 to April 1994. Twenty two children (7 girls, 15 boys) with a mean age of 6-02 years (range 0-83-12-08) were investigated by bronchoscopy for persisting lobar collapse. Ten patients had left lower lobe collapse, nine right middle lobe collapse, five right lower lobe collapse, two right upper lobe collapse, one lingula lobe collapse, and one left lung collapse. Findings at bronchoscopy were as follows: collapse/no infection = 6, collapse/persisting infection = 4, obstruction of bronchi = 7 (three were obstructed by peanuts previously unsuspected, two by mucous plugs), normal = 5. Outcome if bronchoscopy abnormal: resolution of collapse = 6, collapse improved = 2, persistent collapse with evidence of bronchiectasis = 4, lobectomy = 2. Persisting lobar collapse in children should be managed actively to prevent the development of permanent collapse and bronchiectasis. Flexible fiberoptic bronchoscopy is useful in the management of persisting lobar collapse as it can detect causes of bronchial obstruction (including an unsuspected foreign body) and allows suction to clear mucous plugs. We suggest that any child who has lobar collapse that persists after conventional treatment should have a bronchoscopy.

Short term respiratory outcome in infants with chronic lung disease of prematurity (CLD): risk of admission/sudden infant death syndrome (SIDS)

R ILES, AT EDMUNDS The Royal Hospital for Sick Children, Edinburgh Thirty five infants born between 24 and 31 weeks gestation (mean (SD) 27 (1-9) weeks) weighing a mean (SD) of 1010 (71)g at birth were studied between the 39th and 44th week of post-conceptional age. All infants required ventilatory support from birth (mean (SD) 18-7 (3-11) days). Thirty three infants had a chronic oxygen requirement at 36 weeks. Infants were studied using the squeeze technique to measure VmaxFRC and overnight assessment of oxygen saturation (Sao2). Analysis of overnight saturation excluded movement artefact. At time of examination eight infants were on supplementary oxygen (Sao2 was assessed in air for short periods only). No child was on theophylline. The clinical course and the cohort was recorded over the next three months. Nineteen infants were readmitted to hospital with respiratory related illness (76%), twelve with proven respiratory syncytial virus (RSV) infection, eight with a history suggestive of an acute life threatening event (ALTE). One child died unexpectedly, the post-mortem examination was highly suggestive of SIDS showing minimal changes of CLD. For all infants mean Sao2 correlated with risk of admission (t=4.47, p<0.001) and ALTE/SIDS (t=2.8, p<0.001), the variability of the Sao2 correlated with risk of admission (t=2, p<0.05) and ALTE/SIDS (t=3.6, p<0.001). For the non-oxygen dependent infants mean Sao2 correlated with ALTE/SIDS (t=2.3, p<0.05) and ALTE/SIDS with evidence of bronchiectasis (t=3.38, p<0.005). Those infants who were readmitted following RSV infection were graded according to their clinical severity. For this group the mean Sao2 highly correlated with the severity of illness on admission (R=0.77, p<0.005). VmaxFRC did not correlate with clinical outcome.

Influence of maturity at birth on lung volume of preschool children

F GRIFFIN, A GREENHOUSE Paediatric Respiratory Laboratory, Departments of Child Health and Thoracic Medicine, King's College Hospital, London In a prospective follow up study of prematurely born children we demonstrated that, at approximately four years of age, 41% were hyperinflated - that is, they had a functional residual capacity (FRC) greater than 120% of that expected. These infants with chronic lung disease, however, was included, thus it was not possible to determine whether the hyperinflation related to the high proportion of children who were symptomatic or to premature birth per se. We have now therefore matched for height 46 of the prematurely born children (study group) with chil dren born at term who had no respiratory problems (controls). In both groups of children FRC was measured by helium gas dilution. The study group was born at a median gestational age of 29 weeks (range 24-35), 27 had been ventilated in the neonatal period and 15 were symptomatic at four years. The median FRC of the study group (670 ml, range 111-1260 ml) was significantly higher than that of the control group (641 ml, 370-950), p<0.01. No such difference, however, was seen when the asymptomatic prematurely born children were compared with their matched
controls (685 ml, 420–1260 v 711 ml, 400–950). Fifteen of the study group were hyperinflated. The hyperinflated children in the study group did not differ significantly from the rest of their cohort regarding their gestational age (median 29 weeks, 25–35 v 29 weeks, 24–32). A greater proportion of the hyperinflated children, however, were symptomatic (67% v 19%) p<0.01. We conclude hyperinflation in prematurely born children of preschool age does not appear to be influenced by maturity at birth.

Maternal asthma, premature birth and the risk of respiratory morbidity in primary schoolchildren

YJ KELLY, BJ BRABIN, P MILLIGAN, D HEAP, J REID, MG PEARSON School of Tropical Medicine, Alder Hey, Sefton Health and Aintree Chest Centre, Liverpool Maternal asthma and prematurity are known to predispose to the development of later childhood respiratory morbidity. We analysed the relationship of maternal asthma, premature birth and later respiratory outcome in data from parent completed questionnaires collected during two cross sectional community surveys of primary school children, aged 5–11 years, in Merseyside during 1991 and 1993. The numbers of children surveyed were 1872 and 3746 respectively. The proportions of preterm deliveries among asthmatic mothers in the surveys were 23.7% (28/118) in 1991 and 19.1% (61/319) in 1993, compared with 14.0% and 13.0% of non-asthmatic mothers delivering pretermly. Asthmatic mothers are more likely to have a preterm delivery than non-asthmatic mothers (OR 1.49, 95% CI 1.10 to 2.02). Smoking is a separate risk factor for preterm delivery (OR 1.35, 95% CI 1.10 to 1.65). Asthmatic mothers did not have an increased risk of delivering small, growth retarded babies. Maternal asthma, paternal asthma and premature birth, in that order, increased the risk of later childhood respiratory morbidity (OR 3.13, 95% CI 2.36 to 4.16; 2.23, 1.62 to 3.05; 1.40, 1.10 to 1.79). Babies born small but full term are less likely to develop respiratory symptoms in childhood, although this is not statistically significant (OR 0.63, 95% CI 0.28 to 1.41). The effect of maternal asthma on preterm delivery is independent to that of smoking during pregnancy. Maternal asthma is associated with an increased number of premature (but normal size for dates) babies. Prematurity has a lesser effect than genetic factors on predisposing to the later development of respiratory symptoms.

Association of family size with atopic disease

D JARVES, P BURNEY, S CHINN, C LUCZYNSKA, E LAI, R HALL, B HARRISON, J STARK Department of Public Health Medicine, UMDS; Department of Respiratory Medicine, The Ipswich Hospital; Department of Respiratory Medicine, West Norwich Hospital; Department of Respiratory Medicine, Addenbrookes Hospital, Cambridge It has been postulated that atopic disease is inversely associated with family size (Strachan BMJ 1989; 299:1259–60). Data collected as part of the British arm of The European Community Respiratory Health Survey were analysed to determine the association of family size with symptoms suggestive of current asthma and hayfever in a random sample of 1094 young adults living in East Anglia. Of those interviewed 866 (79.1%) agreed to have blood tests for determination of specific IgE to house dust mite (HDM) and timothy grass (Pharmac Diagnostics). The prevalence of symptoms/sensitisation by number of siblings is shown in the table.

<table>
<thead>
<tr>
<th>Number of siblings</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4+</th>
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</thead>
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<tr>
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<td>31</td>
<td>27-8</td>
<td>25-8</td>
<td>32-1</td>
<td>24-5</td>
</tr>
<tr>
<td>Woken by SOB</td>
<td>5-3</td>
<td>11-4</td>
<td>7-8</td>
<td>10-2</td>
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<td>6-0</td>
<td>3-9</td>
<td>2-3</td>
</tr>
<tr>
<td>Hayfever††</td>
<td>33-0</td>
<td>34-3</td>
<td>27-8</td>
<td>28-2</td>
<td>22-2</td>
</tr>
<tr>
<td>Specific IgE to HDM</td>
<td>31-0</td>
<td>26-2</td>
<td>22-4</td>
<td>25-0</td>
<td>22-4</td>
</tr>
<tr>
<td>Grass*</td>
<td>25-4</td>
<td>30-1</td>
<td>23-2</td>
<td>21-8</td>
<td>20-9</td>
</tr>
</tbody>
</table>

*p<0.05, **p<0.01 (χ² for trend significance).
††p<0.05, ‡‡p<0.01 (χ² for trend significance after adjustment for age, sex, smoking and area).

Although there was an association of family size with some of the symptoms, birth order was associated with hayfever only. The prevalence of reported hayfever and sensitisation to grass were negatively associated with family size. The relationship of family size to asthma and sensitisation to house dust mite was less clear. [This work was supported by the National Asthma Campaign.]

Synthesis of IL-8, GM-CSF and TNFα by cultured nasal epithelial cells from atopic rhinitic, atopic non-rhinitic and non-atopic non-rhinitic subjects and the effect of exposure to nitrogen dioxide (NO2)

MA CALDERON, JL DEVALIA, RJ SAPPFORD, A PRIOR, RJ DAVIES Departments of Respiratory Medicine and Allergy, EHT, St Bartholomew’s Hospital, London Recent studies have suggested that airway epithelial cells of atopic and non-atopic individuals may differ in their ability to generate inflammatory mediators. We have cultured nasal epithelial cells (NEC) from biopsy tissues of 10 atopic rhinitic subjects (AR), nine atopic non-rhinitic subjects (ANR), and eight non-atopic non-rhinitic subjects (NNR) and investigated the release of GM-CSF, IL-8 and TNFα and the effect of release of these mediators after exposure for six hours to 400 ppb NO2. Cells were grown as confluent explant cultures on tissue culture inserts and the cytokines released into the medium were estimated by ELISA. NEC from AR released significantly greater (p<0.05) amounts of IL-8, TNFα, and GM-CSF than NEC from ANR and normals under normal culture conditions. IL-8 was generated in greatest quantity and GM-CSF in lowest quantity, irrespective of whether the cells were derived from atopic or non-atopic subjects. Exposure of NEC for six hours to 400 ppb NO2 led to significant release of IL-8 in the normal and the AR group (14-2 and 62 pg IL-8/μg cellular protein, respectively), but not in NEC from the normal group after exposure to NO2. These results suggest that NEC from atopic rhinitic individuals release more GM-CSF, IL-8 and TNFα than NEC from atopic non-rhinitic and normal volunteers and that exposure to NO2 may influence the synthesis of pro-inflammatory cytokines primarily from the NEC of non-atopic non-rhinitic individuals.

Expression of IL-1β and IL-1 receptor antagonist (IL-1ra) in asthmatic bronchial epithelium: effect of steroids

AR SOUSA, SJ LANE, JN KAHKOSTHEN, RN POSTON, TH LEE Departments of Allergy and Respiratory Medicine and Experimental Pathology, UMDS, Guy’s Hospital, London; Department of Respiratory Medicine, Augusta Teaching Hospital, Bochum, Germany Accumulating evidence suggests that the cytokine network is central to the immunopathology of bronchial asthma. Recent evidence for the existence of naturally occurring cytokine antagonists has added to this complexity. In this study we looked at the expression of IL-1β and its naturally occurring receptor antagonist, IL-1ra, in normal and asthmatic bronchial epithelium and have investigated the effect of inhaled beclomethasone dipropionate. Frozen bronchial biopsies from 12 normal and 18 asthmatic individuals were stained with rabbit anti-IL-1β (R&D) and a rabbit anti-IL-1ra (Prof WP Arend, Colorado). Fluorescence-intensity colour image analyses (HIS) were used to quantify the brown immunoperoxidase reaction color present on the bronchial epithelium. There was an increased expression of both IL-1β and IL-1ra in the bronchial epithelial basement membrane, p<0.001 for both. There was a significant correlation between the percentage epithelium stained for IL-β and for IL-ra (r=0.65, p<0.001). Additionally, frozen biopsies from six asthmatic individuals on 1000 μg beclomethasone per day for eight weeks and from six asthmatic individuals on matching placebo, were stained with rabbit anti-IL-1β and anti-IL-1ra antibodies and the immunohistochemical results were quantified by HSI. There was a significant decrease in the percentage change in expression of IL-1β by beclomethasone (p>0.01) but not of IL-1ra (p<0.01). Therefore, IL-1β and IL-1ra are expressed more in the epithelium of asthmatic airways and beclomethasone selectively inhibits IL-1β epithelial expression without affecting IL-1ra significantly.

Increased H2O2 generation by primed human eosinophils following allergen challenge

DJ EVANS, MA LINDSAY, BJ O’CONNOR, PJ BARNES Department of Thoracic Medicine, National Heart and Lung Institute, London; Clinical Studies

Proceedings of the British Thoracic Society

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Unit, Royal Brompton Hospital, London. Activation of inflammatory cells is known to play a part in the pathological changes seen in the asthmatic airway. In this study we have examined the possibility that the eosinophil is activated in asthma using allergen challenge as a model of allergic inflammation. The effect of allergen challenge on the ability of ex vivo human eosinophils to release hydrogen peroxide following stimulation with f Met-Leu-Phe, activation of C5a and complement factor 5a (C5a) was studied in five mild asthmatics. Subjects were venesected immediately before and 24 hours after allergen challenge. All subjects had a late asthmatic response with a mean maximum fall in FEV1 of 29% between 4 and 10 hours after challenge. Freshly separated eosinophils (>95% purity) were incubated in fibrinogen-coated wells (100 μg/ml) for each patient. Cells were serially increasing doses of PAF or C5a. Hydrogen peroxide levels were measured in the supernatant of the cell culture following this incubation. The dose response curves and time course plots showed a dose and time dependent effect for both PAF and C5a. The maximum response for PAF was achieved at a dose of 10-4 M and for C5a, 10-7 M. Allergen challenge resulted in a priming effect on the eosinophils as measured by changes of H2O2 released by the cells. This reached a level of significance for PAF (p<0.02), but not for C5a (p = 0.2). Additionally adhered cells in the absence of stimulation had increased capacity to release H2O2 after allergen challenge. These results demonstrate an in vivo priming of human eosinophils following allergen challenge. This augmented cellular response is likely to be an important factor in the pathogenesis of the late asthmatic response.

Human bronchial epithelial cells express and release RANTES

JL Devalia, JH Wang, RJ Sapsford, RJ Davies Department of Respiratory Medicine and Allergy, St Bartholomew’s Hospital, London. Although several cell types, including T lymphocytes, fibroblasts, endothelial cells and renal tubular epithelial cells, are known to express the chemokine RANTES, to date there is little information on the expression of this cytokine by human bronchial epithelial cells. We cultured human bronchial epithelial cells (HBEC) from surgical tissue in vitro and investigated these for their ability to synthesise and release RANTES, both constitutively and after stimulation with TNFα. Additionally, we have obtained bronchial biopsies from mild asthmatic patients, treated with either inhaled beclomethasone dipropionate (BDP) 500 μg bd or matched placebo for four months and investigated these for the presence of RANTES in the bronchial epithelium, before and after treatment. RANTES in the biopsies was detected by immunocytochemistry and expressed as the percentage of total epithelium staining for RANTES, by use of a colour image analysis system. Measurement of RANTES released from HBEC demonstrated that this cytokine was synthesised constitutively and that this was significantly increased by TNFαs in a dose-dependent manner (range 10-63 fg/μg cellular protein) after 24 hours. Treatment with anti-TNFαs blocked the TNFα-induced release of RANTES from the HBEC. Staining of biopsies demonstrated that RANTES was expressed in the bronchial epithelium of mild asthmatics in vivo and that treatment with BDP led to a significant decrease in the expression of RANTES from 17.12% to 4.22% (p<0.05). These results suggest that human bronchial epithelial cells are capable of synthesising RANTES, both in vitro and in vivo, and may play a part in the pathogenesis of asthma. Furthermore, corticosteroid treatment may influence airway inflammation by modulating the expression and/or release of proinflammatory cytokines in the bronchial epithelium.

Mediators from human bronchial epithelial cells can influence eosinophil and neutrophil chemotaxis in vitro

MM Abdelaziz, JL Devalia, OA Khair, RJ Sapsford, RJ Davies Department of Respiratory Medicine and Allergy, St Bartholomew’s Hospital, London. We have recently demonstrated that human bronchial epithelial cells are capable of releasing proinflammatory cytokines including GM-CSF, IL-8 and TNFαs (Devalia et al. Am J Respir Cell Mol Biol 1993;9:271–9). To investigate the biological relevance of the release of these cytokines we have collected 24 hours conditioned medium (CM) from confluent explant cultures of human bronchial epithelial cells. The effects of this medium on the chemotactic response of preparations of eosinophils and neutrophils isolated from normal individuals. Chemotaxis was assessed according to the Boyden technique and results were expressed as the mean number of cells in 10 random high power fields. CM significantly increased eosinophil chemotaxis (median 11-4, range 9-1-12-3; p = 0-005) compared with medium 199 (negative control, median 4-1, range 3-4–7-0), but not when compared with 10 μM PAF (positive control, median 20-0, range 13-0–26-5). Similarly, CM also significantly increased neutrophil chemotaxis (median 30-4, range 19-4–33-3; p<0-01) compared with medium 199 (median 17-2, range 13-4–20-9). A positive control with 10-6 M f Met-Leu-Phe (positive control, median 64-7, range 49-0–132-0). These results suggest that human bronchial epithelial cells produce inflammatory mediators which can influence the activity of eosinophils and neutrophils and consequently may play an important part in the aetiology of airways disease.

Accident and emergency attendance for asthma: an opportunity for intervention?

PJ Vickery, RJ Miconough, DPS Spence, CRK Hind Royal Liverpool University Hospital Trust (RLUHT), Liverpool. The attendance of an asthmatic patient at an Accident and Emergency (A&E) department is a failure of treatment. Patients attending A&E may be either admitted to the hospital under the care of physicians, admitted to the short stay observation ward (SSOW) in A&E. The care of those admitted to hospital under physicians has been audited nationally (Pearson et al. Thorax 1992). However, patients whose care has been A&E based have not been so well examined. Since October 1993 an asthma liaison nurse (ALN) has been attached to the A&E department of the RLUHT and is responsible for assessing patients in the SSOW and advising appropriately. In previous case-control studies examining the association between inhaled β agonist use (positive “index event”) and emergency department attendance, only 25% had attended for asthma review in the previous year. 64% of patients had sought help for their asthma since increasing symptoms had begun. Despite the fact that 72% of the GP practices of these patients were approved for asthma chronic disease management programmes by the Family Health Services Authority, only 25% had attended for asthma review in the previous year. 64% of patients had sought help for their asthma since increasing symptoms had begun. Despite the fact that 72% of the GP practices of these patients were approved for asthma chronic disease management programmes by the Family Health Services Authority, only 25% had attended for asthma review in the previous year. 64% of patients had sought help for their asthma since increasing symptoms had begun. Despite the fact that 72% of the GP practices of these patients were approved for asthma chronic disease management programmes by the Family Health Services Authority, only 25% had attended for asthma review in the previous year. 64% of patients had sought help for their asthma since increasing symptoms had begun. Despite the fact that 72% of the GP practices of these patients were approved for asthma chronic disease management programmes by the Family Health Services Authority, only 25% had attended for asthma review in the previous year. 64% of patients had sought help for their asthma since increasing symptoms had begun. Despite the fact that 72% of the GP practices of these patients were approved for asthma chronic disease management programmes by the Family Health Services Authority, only 25% had attended for asthma review in the previous year. 64% of patients had sought help for their asthma since increasing symptoms had begun.

Risk of severe life threatening asthma (SLTA) and death and type of prescribed β agonist: an example of confounding by severity

JE Garrett, S Lanes, J Kolbe, HH Rea Respiratory Services, Green Lane Hospital, Auckling, New Zealand; New England Epidemiology Institute, Newton Lower Falls, Massachusetts, USA. In previous case-control studies examining the association between inhaled β agonists in death or near death, the potential for confounding by disease severity has prompted confirmation of the causal association between asthma drugs and death or SLTA. The aims of this study were (1) to assess the relationship between inhaled β agonists (specifically fenoterol) and other asthma drugs and the risk of asthma death or SLTA, and (2) to examine the influence of confounding by severity by controlling for a variety of risk factors not reliably collected in previous studies. A retrospective, dynamic cohort study was performed on patients aged 13–52 years following a hospital visit (ER visit or admission) between May 1986 and December 1987. Following the “index event” each subject contributed person time to the analysis until death or April 1989, allowing incidence rates for SLTA or death to be calculated. The rates of each outcome in relationship to each drug were calculated, as were incident rate ratios (RRs) for users to non-users. For each type of drug adjusted RRs were calculated, controlling for four risk factors.
factors which were independently associated with an increased risk of SLTA (race, previous hospital admissions, usual asthma symptoms, and severity of previous attack). Six hundred and twenty-four patients were recruited subsequent to which there were 14 asthma deaths, 94 SLTAs, 333 hospital admissions, and 520 ER visits. Crude RRs for SLTAs and death were strikingly similar and were increased for all asthma medications. After controlling for the independent risk factors, most of the crude RRs reduced to a non-significant association. Controlling for hospitalisation in the past year and any oral corticosteroid use reduced the RR for inhaled fenoterol when compared with inhaled salbutamol from 2.02 to 1.52 (CI 1.02 to 2.28). When the four independent risk factors were controlled for, the RR for fenoterol reduced further to 1.11 (CI 0.57 to 2.20). These data support the notion that most asthma medications are related to factors resulting in an attack of asthma and severity of asthma. Users of fenoterol had a higher incidence of life-threatening asthma than users of salbutamol. However, after adjusting for differences in baseline severity, users of fenoterol and of salbutamol had similar risks of life threatening asthma. [Supported by the Asthma Foundation of New Zealand.]

**Beta agonist type and outcome following severe life threatening attack of asthma (SLTA)**

J KOLBE, S LANCES, JE GARRETT, H REA Respiratory Services, Green Lane Hospital, Auckland, New Zealand; The Canterbury Research Group, Christchurch, New Zealand

Confounding by severity remains a tenable explanation for the reported association between prescription of the β agonist fenoterol and asthma death or SLTA. Previous SLTA is associated with a markedly increased risk of asthma death and further SLTA. The aim of this study was (1) to assess the relationship between inhaled β agonists, specifically fenoterol, and the risk of asthma death or SLTA, in a group of subjects at recognised high risk, and (2) to examine the influence of confounding by severity by controlling for a variety of risk factors. A retrospective dynamic cohort study was performed on patients aged 15-50 years following an intensive care admission for SLTA (median 4 (IQR 3), Paco2, 11.0 (5-5) kPa on admission). Each subject contributed person time to the analysis until death or April 1989, allowing incidence rates for SLTA or death to be calculated. The rates of each outcome were calculated as were the incident rate ratios (RRs) for fenoterol to salbutamol users. Adjusted RRs were calculated by controlling for risk factors associated with an increased risk of SLTA in non-fenoterol users. Three hundred and seventy-eight patients formed the cohort. There were 13 deaths, 62 subsequent SLTAs and 345 subsequent hospital admissions. Prescription of fenoterol was associated with an increased risk of SLTA (crude RR = 1.72). Controlling for sociodemographic factors was associated with an adjusted RR of 1.44 (95% CI 1.05 to 1.97). After additional adjustment for multiple risk factors (age, prior SLTA, number of types of inhaled β agonist prescribed), the adjusted RR was 1.02 (95% CI 0.57 to 1.83). We conclude that, in this high risk group, the increased risk of SLTA associated with fenoterol prescription seems best explained by confounding. [Supported by the Asthma Foundation of NZ and the NZ Lottery Board.]

**Comparison of nebulisation therapy combining ipratropium bromide (0.5 mg) plus salbutamol sulphate (3.0 mg) with salbutamol sulphate (3.0 mg) alone in acute asthma**

JE GARRETT, I TOWN for the New Zealand Combinitive Study Group Respiratory Services, Green Lane Hospital, Auckland, New Zealand; The Canterbury Research Group, Christchurch, New Zealand

There is uncertainty about the benefit of routine addition of ipratropium bromide (IB) to β agonist therapy in acute asthma. Patients with more severe acute asthma appear to benefit more from combination therapy. We aimed to evaluate (1) whether a combination of inhaled IB (0.5 mg) plus salbutamol sulphate (SS) (3.0 mg) confers significant additional bronchodilating effects over nebulised SS (3-0 mg) in patients with acute bronchial asthma, and (2) whether adjusting for the effects of known prognostic indicators of outcome (baseline PEV1, age, duration of asthma attack and medicines taken in the six hours prior to presentation) influences any of the additional effect seen with IB. A double blind, two centre, randomised, parallel group, single dose study of 338 asthmatics aged 18-55 years attending an emergency department (ED) with acute asthma. The primary end point was FBV measured at 45 and 90 minutes. The mean (SE) absolute difference in FEV1 with IB plus SS compared with SS at 45 minutes was 93 (44) ml (p<0.05) and at 90 minutes 113 (48) ml (p<0.05). Treatment response was predicted by baseline FEV1, (p<0.001), time between onset of asthma attack and start of nebulisation (p<0.05), age (p<0.05), and inhaled β agonist (p<0.0001) and inhaled anticholinergics (p<0.01) used in the previous six hours. The added advantage of combination therapy response over β agonist response was apparent in all subgroups defined by these prognostic factors. Those subjects who had taken no asthma medication in the six hours prior to ED attendance exhibited the greatest response to combination therapy. Thus, there was a significantly greater response at both 45 and 90 minutes with the combination therapy. Patients with more severe asthma at presentation did not exhibit greater benefit from the addition of IB. Patients who exhibited most benefit from IB were those who had consumed least bronchodilator prior to presentation and who were presumably on the lower part of the dose response curve. [Supported by Boehringer Ingelheim.]

**Peak flow criteria for hospital admission or discharge in acute asthma**

I DAVIES, I RYLAND, M ZAHER, E KADZOMBE, MG PEARSON Accident and Emergency Department, Fazakerley Hospital, Liverpool

The asthma guidelines recommend that patients with acute severe asthma with a peak flow <50% of best or predicted value should be admitted. We studied retrospectively 100 consecutive patients (median age 33, range 5-68 years, 56% female) presenting with acute asthma requiring hospital management. Sixty seven patients (group 1) were admitted, 31 (group 2) were discharged. Two were transferred to a paediatric hospital. Those admitted were older (median 66 ± 23 years), included more smokers (57% vs 29%), had more previous admissions (56% vs 32%) and more on oral steroids (36% vs 10%) (all p<0.02). Use of inhaled bronchodilators (82% vs 80%) and inhaled steroids (69% vs 61%) was similar. All 18 patients with previous ITU admissions and all 16 with home nebulisers were admitted. peak Flows on arrival were recorded in 88 patients; nine of those without initial peak flows were subsequently admitted for SLTA. Peak flow on arrival were lower (p<0.01) in those admitted (median 150/min, range unrecordable-410) than in those discharged (median 330/min, range unrecordable-500) with a marked overlap between the groups (figure). More patients admitted had a tachycardia >110/min (33% vs 17%) and were tachypnoeic (rate >25) (55% vs 41%). Of those sent home 5/31 (16%) adults had a peak flow <200/min on arrival. Four were female and two also had a tachycardia >110/min and respiratory rate >25. Working to guidelines, these patients should have been admitted, which would result in 5/67 (7%) more admissions. However none died or were readmitted within two months. Whether the peak flow thresholds in the guidelines are adequate requires further evaluation.

**Follow up of asthma patients in hospital outpatient clinics**

I RYLAND, MG PEARSON on behalf of the National Asthma Task Force and 30 PARTICIPATING PHYSICIANS Accident and Emergency Centre, Liverpool, England

There are 34 reported claims that hospital doctors follow up too many asthma patients in their clinics. The Asthma Task Force has reviewed 21 adult and nine paediatric hospitals collecting data on 894 adult and 401 paediatric patients. Median age of adults was 48 (13-93) years and of children 6 (0.5-17) years. 59% of adults and 39% of children were female, and 65% adults and 62% children had...
had hospital admissions. Of the adults, 14% were on oral steroids, 24% on high dose inhaled (≥2 other therapy, 38% on low dose steroids (≥2 other therapy), and 24% on no anti-inflammatory treatment. For children 2% were on oral steroids, 20% on high dose inhaled steroids, 25% low dose steroids, 10% cromoglycate, and 45% on no anti-inflammatory treatment. Main reasons given for requiring follow up were asthma severity 43/740 (58%), concomitant disease 81/740 (11%), and special patients (e.g. staff) 81/740 (11%). For children these same reasons applied to 185/298 (68%), 28/298 (9%), and 19/298 (6%) respectively. 112/740 (15%) adults and 22/298 (7%) children were discharged at that appointment with discharge being anticipated within six months for a further 17% and 16% respectively. Of 353/740 (48%) of adults and 118/298 (40%) of children in whom long term follow up was planned, these three reasons accounted for 90% of cases. 240/312 (77%) adults and 83/131 (63%) children being followed long term were on high dose steroids or more. When patients were asked who they would wish to be followed up by (if their asthma was well controlled), 338/708 (48%) said GP alone, 290/708 (41%) hospital and GP and only 80/708 (11%) hospital alone. This was independent of asthma severity as judged by treatment stage. Equi- valent figures for paediatrics were 114/295 (39%), 125/295 (42%) and 118/298 (39%). Many patients would like to be followed up by GP alone. Hospital clinics seem to be discharging most mild asthma patients so that the majority of asthma patients being followed up long term do have severe disease. [Supported by a grant from the Department of Health.]

Increased nitric oxide in exhaled air of normal subjects with upper respiratory tract infection

SA KHARITONOV, DH YATES, PJ BARNES Department of Thoracic Medicine, National Heart and Lung Institute, London Nitric oxide (NO) is produced by a variety of cells within the lower respiratory tract, including inflammatory and epithelial cells, and is increased in inflammatory lung disorders. To determine whether exhaled NO is increased by upper respiratory tract infection (URI) we measured exhaled NO in a total of 18 (11 men) normal subjects (mean age 33-7 (1-05) years) during and after URI, and compared values with age and sex-matched controls. All subjects with URI had documented symptoms, such as generalised or frontal headache, myalgia, rhinorrhea, sneezing, congestion/blocked nose, cough and chest pain accentuated by coughing (14 subjects) for at least two days prior to the investigation. Lung function parameters and exhaled nitric oxide (NO) were investigated in all subjects on two occasions: 1–2 days after the beginning of URI, and three weeks thereafter. Exhaled NO was measured on a chemiluminescence analyser (Dasibi Environmental Corporation, Glendale, California, USA). Subjects were asked to produce a slow vital capacity manoeuvre over 30–45 seconds into wide bore Teflon tubing. NO was sampled continuously. The control group had been investigated earlier in the year and were used as a reference group regarding exhaled NO levels. Mean peak exhaled NO concentration was 87-7 (2-67) ppb in this group of 72 non-smoking controls (43 men mean age 36-1 (2-4) years). In subjects with URI the peak was significantly higher (314-6 (64) ppb, p<0-001) during the acute phase of URI, whereas peak NO values after recovery fell to 87-4 (8-93) ppb, not significantly different from the normal controls. No change was observed in lung function. These data suggest that exhaled NO reflects both upper and lower respiratory tract inflammation and may have clinical utility in monitoring cyto- kine-mediated inflammation.

Nitric oxide production by human bronchoalveolar lavage cells in culture

PL LEUNG, K ANDERSON, C MCSHARRY, IB MCINNES, NC THOMSON, PJ LIWE Department of Immunology and Respiratory Medicine, Western Infirmary, Glasgow There is increasing evidence from both animal and human studies suggesting that nitric oxide (NO) produced by inducible nitric oxide synthase (iNOS) may play an important role in the pathophysiology of certain lung diseases. We sought to establish whether cells obtained by lavage during diagnostic bronchoscopy can produce NO. Bronchoalveolar lavage (BAL) from 23 patients with mainly airway disease but also included patients (10 endobronchial cancer, three infection, six haemoptysis with negative bronchoscopy, age 29–84) were cultured in RPMI/5%FCS for 72–96 hours and NO production was measured by Griess reaction. Expression of the iNOS gene was assessed by reverse transcription polymerase chain reaction (RT/PCR) using oligonucleotide primers corresponding to unique human iNOS sequence. In 15 of 19 samples there was no evidence of NO production after 72–96 hours in culture. The four samples (three bronchial carcinoma) which produced NO did so at low concentration (range 7–13 nmol/l 106 cells/ml) and was not increased even after stimulation with LPS and SHP; this NO production was inhibited in the presence of 5 mm L-NMMA. iNOS mRNA was detected in four of 10 patients and in one patient NO was detected but PCR was negative. The majority of BAL samples did not produce NO either spontaneously or after stimulation. The few positive were mainly from subjects with malignant airways disease.

1-Arginine increases exhaled nitric oxide in normal subjects

SA KHARITONOV, G LUBEK, B LUBEK, M HJELM, PJ BARNES Department of Thoracic Medicine, National Heart and Lung Institute, London; University Hospital, Vienna, Austria; Hospital for Sick Children, London We have investigated the effect of orally administered L-arginine (0-05, 0-1, 0-2 g/kg) compared with matched placebo on the concentration of nitric oxide (NO) in the exhaled air in 23 normal individuals. Administration of the dose of L-arginine (or placebo) was designated time 0 and exhaled NO was measured prior to medication, then every 15 minutes up to five hours after medication. A blood sample was taken for measurement of nitrate and arginine prior to dosing then two hours after dosing. Heart rate, supine blood pressure, and FEV1, were measured hourly throughout the study. The doses of L-arginine were 0-1 g/kg (n = 7) and 0-2 g/kg (n = 4), but not 0-05 g/kg (n = 4), induced a significant increase in exhaled NO two hours after administration (baseline 64 (7-26) ppb; two hours 97 (4-88) ppb, 52 (6-59) ppb, and 156 (19-3) ppb, respectively) which appeared to be dose-related. This was paralleled by a dose-related increase in plasma nitrate and arginine concentrations two hours after administration. There was no effect of placebo (five subjects) on the levels of exhaled NO over the same time period, nor on plasma nitrate or arginine concentrations. The increase in exhaled NO was maximal at two hours and then declined by 3–4 hours to baseline levels. Calculation of the area under the curve by planimetry demonstrated a dose-related increase (p<0-01) in exhaled NO (10-2 (6-6) arbitrary units at 0-05 g/kg, 41-1 (8-8) units at 0-1 g/kg, and 114-5 (24-0) units at 0-2 g/kg). There was no significant effect on heart rate, blood pressure, or FEV1, over the period of measurement. The results suggest that an increase in the amount of substrate for NO synthesis can increase the formation of endogenous NO.

Downregulation of the constitutive nitric oxide synthase (NOS) gene expression in rats treated with lipopolysaccharide in vivo

SF LIU, IM ADCOCK, PJ BARNES, TW EVANS Unit of Intensive Care, Department of Thoracic Medicine, National Heart and Lung Institute, London Bacterial endotoxin has been reported to be both stimulatory and inhibitory on nitric oxide release. One explanation for this discrepancy is that endotoxin may affect the constitutive and inducible NOS (iNOS) gene differently. We have reported that lipopolysaccharide (LPS) treatment of rat in vivo causes a widespread tissue expression of the iNOS gene. We now study the effects of LPS and dexamethasone (Dexa) treatment in vivo on the endothelial type (eNOS) and brain (nNOS) mRNA expression in the rat. Glyceraldehyde phosphate dehydrogenase (GAPDH) gene was also studied in parallel as an internal standard. Wistar rats were treated with either saline (1 ml/kg ip, control group) or LPS (15 mg/kg ip, LPS group) for four hours. For LPS + Dexa group, Dexa (3 mg/kg ip) was injected 40 minutes prior to the administration of LPS. Lung, heart, brain and testis were dissected from these animals. Total and mRNA were isolated using commercial kit. Northern blot analysis was performed according to standard methods. The eNOS and nNOS mRNA level were quantified using densitometry and expressed as eNOS/GAPDH or nNOS/GAPDH ratio. Neither LPS nor Dexa had any effect on the GAPDH mRNA levels. The eNOS/GAPDH ratios for control, LPS, and LPS + Dexa groups, respectively, were 0-31 (0-07), 0-11 (0-04), and 0-08 (0-02) (p<0-03 compared with control, n = 6) in the lung, and 0-16 (0-03), 0-04 (0-02), and 0-06 (0-02) (p<0-03 compared with control, n = 6) in heart. Similar results were obtained from aorta. The bNOS/GAPDH ratio was 0-42 (0-15), 0-10 (0-02), and 0-06 (0-01) (p<0-03 compared with control) for control, Endo, and Endo + Dexa group respectively. These results demonstrate that LPS treatment in vivo downregulates the two isoforms of con-
Hypoxic pulmonary vasoconstriction/dilation is modified by different preconditions

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In vitro, hypoxia causes both constriction and dilatation. Small vessels were mounted in a Cambusmyograph in physiological saline in the isometric mode, gassed with O₂ or N₂ + 5% CO₂. After preconstriction with 4 μg PGF₂α, hypoxia caused a four-phase response: (1) dilatation; (2) constriction; (3) dilation; and (4) a slow small constriction. This response was repeatable. The response varied with the level of preconstriction and with different agonists. In 20 vessels (five rats), at lower levels of preconstriction with PGF₁α, hypoxia caused small prolonged contractions only; both dilator and constrictor responses increased with the level of preconstriction. During KCl preconstriction, dilatation and constriction were less than after PGF₂α (phase 1, 0-017 (0-01) v 0-21 (0-02) mm/mm; phase 2, 0-33 (0-097) v 2-19 (0-21) mm/mm, both p<0.001), even though the preconstriction level was similar (PGF₂α, 0-99 mm Hg; PGF₁α, 0-91 mm Hg). In the maximal contraction of eight pulmonary vessels, (four rats) to KCl was unchanged (3-35 (0-91) mm Hg/v 3-71 (0-99) mm Hg, p=0-2661) compared with 38°C, but PGF₂α, maximal contraction was reduced (1-48 (0-32) mm Hg/v 3-08 (0-61) mm Hg, p<0.001). Acetylcholine (ACH) relaxation was higher at 25°C (104 (56)%) compared with 38°C (12 (10)%). PGF₂α, preconstriction (p<0.001). The hypoxic response was tested at 25°C in eight vessels (four rats) preconstricted with PGF₁α. The magnitude of contraction by hypoxia compared with 38°C was unchanged. However, at 25°C, the onset of hypoxic constriction was delayed (2-83 (0-6) min at 38°C v 6-87 (1-24) min at 25°C, p<0.001). The contraction was maintained for a longer time (7-83 (1-64) min at 38°C v 24-62 (4-34) min at 25°C, p<0.001), and dilatation was absent.

Energy state, intracellular pH and tension responses in pulmonary artery rings during hypoxia and absence of substrate

RM LEACH, DW SHEEHAN, VP CHACKO, JT SYLVESTER Department of Intensive Care, St Thomas’ Hospital, London

Reduction in energy state may trigger hypoxic pulmonary vasoconstriction and is associated with hypoxic systemic vasorelaxation. 13P nuclear magnetic resonance spectroscopy (NMR) was used to measure phosphocreatine (PCr), ATP and intracellular pH (pHi) in porcine intrapulmonary artery rings. Paired rings were stretched to 4 g tension in standard organ baths. One ring from each pair was transferred to a plastic frame at in vitro temperature (102mmHg) placed in a sample tube within the bore of an 11-8 Tesla MSL 500 NMR spectrometer, and superfused with phosphate-free Krebs-bicarbonate solution (37°C, pH=7-4) containing phenylephrine (10⁻⁴ M). NMR spectra were obtained from free induction decays collected over 15 minutes. The other ring remained in the organ bath to allow measurements of tension and pH to be subjected to identical conditions. Experiments consisted of control, experimental and recovery periods each two hours in length. In the control and recovery periods the perfusate contained glucose (10 mM) and was gassed with 93% O₂. In the experimental period the perfusate was either gassed with 0% O₂ (Po<20 torr) or glucose was removed (sucrose substitution) or both. During hypoxia PCr transiently fell to 37 (14%) of baseline returning to 90 (10%) at two hours. Initially, pHi also fell from the control value of 7-24 (0-02) to 7-08 (0-03) recovering to 7-18 (0-02) after two hours hypoxia. ATP was unaffected. These changes correlated with initial vasoconstriction followed by vasodilation. Removal of glucose during hypoxia prevented vasoconstriction, inhibited the recovery of PCr and pHi, and resulted in a fall in ATP. Absence of glucose without hypoxia had no effect on tension, energy state, or pHi. The results suggest a link between energy state, pH and pulmonary vasoconstrictor tone. [RML supported by MRC.]

Endothelin receptors in rat pulmonary arteries and arterioles: effect of pulmonary hypertension

KM MCCULLOCH, M BARD, MR MACLEAN Division of Neuroscience and Biomedical Systems, IBLIS, West Medical Building, Glasgow University, Glasgow

In adult rats vasoconstrictor responses to endothelin-1 (ET-1)
Cyclic GMP levels are raised in endothelium-denuded rat pulmonary artery after LPS treatment

**MJD GRIFFITHS, M MESSENT, NP CURZEN, TW EVANS** National Heart and Lung Institute, London Nitric oxide (NO) acts on vascular smooth muscle through activation of soluble guanylyl cyclase as a mediator of guanosine 3',5'-cyclic monophosphate (cGMP) (Circ Res 1983;52: 352-7). We have demonstrated pharmacologically that endotoxin (LPS) injection induces NO synthase (NOS) in rat pulmonary vascular smooth muscle (PVSM) (Thorax 1993;48:467). Biochemical confirmation of NOS induction in PVSM requires the demonstration of raised levels of cGMP in vessels from LPS-treated rats that could be decreased by an NO inhibitor, eg. N^6^-monomethyl-L-arginine (LNM). Male Wistar rats were treated with LPS (20 mg/kg ip) or vehicle (1 ml NaCl, ip) four hours before sacrifice. Main pulmonary artery rings were suspended in organ baths containing oxygenated Krebs solution. Their endothelium was removed by abrasion, confirmed by failure to relax to acetylcholine after application of a phenoxybenzamine (PE) dose-response curve. After equilibration, vessels were contracted by their EC50 PE and then by LN (50 μM or 1 mM) or vehicle until a plateau tension was reached when the rings were snap frozen and stored at -80°C prior to radioimmunoassay for cGMP (Adv Cyclic Nucleot Res 1979;10:1-33). Results shown in the table are the mean (SE)

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<td>Tm (%)</td>
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*p<0.05, **p<0.01 LN [0] v LN [50 μM or 1 μM].

LN-induced tension (Tm) expressed as % contraction caused by the PE EC50 and the cGMP level (fmol/mg, n = 8). cGMP levels were increased following LPS treatment and diminished, in a dose-dependent manner by NOS inhibition, suggesting that NO activity was increased following NOS induction in PVSM by LPS. [MG is a Wellcome Clinical Research Fellow.]

Responses to 5HT and sumatriptan in control and pulmonary hypertensive rats

**MR MCLEAN, M BAIRD, KM MCCULLOCH** CRI Laboratories, Division of Neuroscience and Biomedical Systems, IBLS, West Medical Building, Glasgow University, Glasgow We have previously shown that contractile 5HT_1D receptor-mediated responses to sumatriptan exist in bovine and human pulmonary arteries and that in these vessels, unlike in the systemic circulation, the 5HT_1D receptor-mediated responses are sensitive to nitric oxide (NO) donors. We have now examined the effect of 5HT on isolated rat pulmonary arteries treated with LPS (a potent inducer of NO synthase (NOS) activity) and a specific inhibitor of NO. We have used 5HT to examine the changes in pulmonary arterial contractility in response to NO donors and to determine the role of NO in the regulation of pulmonary arterial tone in vivo.

**Vasodilator efficacy of anitrosothiol SNAP and SNP in normal and chronic hypoxic rat lung**

**CELA EMEY** Department of Medicine, University of Sheffield, Sheffield s-Nitrosothiols, which liberate nitric oxide, may mediate

Endothelin-1 induced contraction in human pulmonary arteries: mediated by ETA receptors in large arteries but not ETA/non-ET receptors in resistance arteries

**KM MCCULLOCH, M BAIRD, MR MCLEAN** Division of Neuroscience and Biomedical Systems, IBLS, West Medical Building, Glasgow University, Glasgow In rats vasocostrector responses to endothelin-1 (ET-1) are mediated via the ETA receptor in pulmonary arteries, but not through both ET-A and ET-B receptors in pulmonary resistance arteries (PRAs) (MacLean et al. J Cardiovasc Pharmacol 1994:23:838-45). Here we investigated the effect of pulmonary hypertension (PH) induced by exposure to hypoxia in young SPF rats. Chronic hypoxic (CH) rats (8-30 days old) exposed to 10% O2 for two weeks and age matched controls were studied. The following pulmonary arteries (PAs) were set up under 1-5 g tension in 5 ml organ baths. The PAs (150 μm id) were set up on a wire myograph under 130-40 mg tension. All vessels were bubbled with 16% O2/6% CO2. Response curves were obtained to ET-1 (0-1, CH rats: 0-1, n=10), ETA agonist sarafotoxin S6c (SSS6c) (n=5-10 rings in 6-10 rats). Control rat PAs contracted only to ET-1 (log M EC50 -8-4 (0-2)) and these responses were inhibited by FR93931, indicating contractions via the ETB receptor. CH rats contracted to both ET-A (which was more potent than in controls, p<0.05, Student's unpaired t test, log M EC50 -8-9 (0-1)) and SSS6c (log M EC50 -7-7 (0-2), maximum response: 30% compared to ET-1) indicating contraction through the ETB receptor and an ETB-like receptor. FR-sensitive responses to ET-1 were seen in both control and CH PAs (log M EC50 controls -8-1 (0-1), CH rats: -8-1 (0-1)). SSS6c also induced potent contractions in the PRAs (log M EC50 controls -9-2 (0-1), CH rats: -9-9 (0-1)). The maximum response to ET-1 was increased in the CH rats by 50% whilst the maximum response to SSS6c was unchanged. This study shows that ET-1 contraction is mediated only by ETB receptors in these rat PAs but that ETB receptors play a major role in PRAs. In PAs, PH induced an increase in the ETB-mediated response. CH rats that were exposed to LPS (1 μM) for two weeks and their age matched controls were studied. PH was confirmed by multiplying right ventricular/total ventricular weight ratio which was 60% greater in the CH rats. The right branch pulmonary arteries (PAs) were set up as rings, suspended in organ baths containing oxygenated Krebs solution and were either a) precontracted with ET-A or b) precontracted with ET-B and then exposed to 16% O2/6% CO2. Cumulative concentration response curves (CCRC) were constructed to both 5HT and sumatriptan. In the latter experiments, a maximum concentration of 5HT was administered at the end of the CCRC to sumatriptan. Responses to 5HT were seen in the presence of 5HT (n=6) in control PAs and were 7-6 (0-1) in CH rat PAs (p<0.001 compared with controls; Student's paired t test). In the control PAs there was no response to sumatriptan whereas the CH rat PAs contracted to sumatriptan which gave a maximum response 40%-50% of that to 5HT. The log M EC50 value for sumatriptan was -6-0 (0-5) (n=6) and therefore equipotent with 5HT in the control PAs. In conclusion, only are 5HT, ET-A induced contractile responses important in PAs, but this study suggests that they are potentiated in pulmonary hypertension and may account for the increased response to 5HT observed in CH rats. [This work was funded by the MRC.]
endothelial-dependent vasodilatation. The effect of s-nitroso acetyl penicillamine (SNAP) on pulmonary vascular tone (low during normoxia or raised by either chronic hypoxia (10% O_2 three weeks) or during acute hypoxic vasoconstriction (HPV)) was tested in isolated rat lungs (IPL) and compared with the effect of SNP, sodium nitroprusside, a non-endothelial dependent dilator. Lungs were perfused in situ with homologous blood at a constant flow of 20 ml/min; and ventilated with a mixture of N_2O (20% O_2, 5% CO_2) and 95% O_2, equivalent to 0.3 ml bolus doses of SNAP or SNP (0-01, 0-1, 1, 10, 100 μg) on pulmonary artery pressure (Ppa) was studied in both normal (C) and chronically hypoxic (CH) IPLs. During normoxia both SNAP and SNP caused small falls in Ppa, being more obvious in CH (range 0-5-3.5 mm Hg). During hypoxia SNAP and SNP caused dose-dependent falls in Ppa, expressed as % fall in HPV, similar in C and CH rats. However, SNP was 10 times more potent than SNAP (100% fall in HPV with 100 μg SNAP v 10 μg SNP). The nitric oxide synthase inhibitor, t-NAM, had no consistent effect on the vasodilatation. Thus, the nitrosothiol, SNAP, causes profound vasodilatation in the rat lung, equipotent in C and CH, although potency was less than SNP. Neither were consistently affected by blockade of endogenous NO release. Inhaled nitrosothiols may provide an alternative treatment to nitric oxide in pulmonary hypertension.

Nebulised prostacyclin used as a selective pulmonary vasodilator in acute lung injury

J SHEPHARD, SJ BRETT, B KEogh, TW EVANS Unit of Critical Care, Royal Brompton Hospital, London Prostacyclin (PGI_2) has been widely used intravenously as a vasodilator in a variety of pathophysiological circumstances, including sepsis and pulmonary hypertension. There are theoretical and practical advantages to using a selective and locally administered vasodilator in the management of pulmonary hypertension and hypoxia in patients with ARDS. We describe the response to nebulised PGI_2 of four patients with ARDS secondary to a variety of pathologies. Prior to PGI_2 administration they were managed on pressure controlled, inverse ratio ventilation, which was continued during therapy. Prostacyclin was administered via a gas-driven nebuliser attached to the distal portion of the inspiratory limb of the ventilator circuit. Baseline measurements of pulmonary (PVR), systemic (SVR), vascular resistance and PaO_2/FiO_2 ratio (P/F) were made. The response to increasing doses of PGI_2 were then assessed. The results are shown in the table as % changes from baseline. The minimum effective dose ranged from 4 to 32 ng/kg/min. These data show that PGI_2 may be successfully administered via nebuliser to patients with ARDS. Its effects were to reduce intrapulmonary shunt fraction and PVR without significantly changing SVR. The wide variation in dose response between patients shows the value of performing a dose response for each patient.

Prediction of pulmonary hypertension in patients with systemic sclerosis and fibrosing alveolitis

JB CAILES, CM BLACK, RM DUBoIS Royal Brompton Hospital, London Systemic sclerosis (SSc) is often associated with fibrosing alveolitis (FASSc). Pulmonary hypertension (PHT) is the major cause of mortality in this condition. Precise estimation of pulmonary artery pressure requires the invasive procedure of right heart catheterisation. Doppler echocardiography (DE), however, is both sensitive and specific for the diagnosis of PHT in SSc but this test is not routinely available in all hospitals. We aimed to identify the demographic, disease and lung function factors which would most reliably predict risk of PHT in patients with FASSc. One hundred and thirty patients with FASSc prospectively underwent Doppler echocardiography. In 78% of Doppler green the pulmonary artery systolic pressure was greater than 30 mm Hg (29 patients). Exclusion of PHT required demonstration of a structurally normal right ventricle and a normal time interval between closure of the pulmonary valve and start of tricuspid flow on Doppler. Patients with evidence of pericardial disease or reduced left ventricular function were excluded leaving 113 patients for analysis. Demographic data (age, sex, smoking history), disease variables (duration of pulmonary symptoms, and type of SSc), and full lung function were collected at the same time as the DE. Multiple linear logistic regression was used to model the relationship between these variables and the presence of PHT as determined by DE. After matching for other indices transfer factor expressed as the percentage of predicted normal, corrected for Hb (TLco) and the alveolar to arterial oxygen gradient (Aa) were the only significant independent variables for the prediction of PHT (p values of 0.0014 and 0.022, respectively). A logistic regression equation with presence or absence of PHT as the dependent variable and independent variables of age, sex, smoking history (ever v never smokers), SSc type, duration of pulmonary symptoms, TLco and Aa, correctly designated 91-2% of cases. Twenty one of 29 patients with PHT were correctly identified (72-4%), whilst 82 of 84 patients without PHT were correctly assigned (98-8%). Using TLco and Aa alone as the independent variables, 89-4% of cases were correctly assigned using the equation:

\[
\text{probability} = 1/1 + \exp\left(-1.187 - 0.0867 \times \text{TLco} + 0.604 - 3.032 \times \text{Aa}\right)
\]

with a probability value of 0.5 representing PHT. We conclude that in patients with FASSc, both TLco and Aa gradient for oxygen are independently predictive of PHT. TLco and arterial blood gases should be performed as part of routine assessment of patients with FASSc, and can predict patients with a high likelihood of having PHT. [Supported by a grant from the Raynauds and Scleroderma Association UK.]

Alterations in CT parenchymal density gradients in lone pulmonary hypertension complicating systemic sclerosis

JB CAILES, RM DUBoIS, D M HANSELL Royal Brompton Hospital, London Lone pulmonary hypertension (PHT) occurring in the absence of fibrosing alveolitis (FA) is a rare but well recognised complication of systemic sclerosis (SSc) which carries a poor survival prognosis. Histologically there is fibrointimal proliferation and obliteration of the pulmonary microvasculature. Disturbances of the pulmonary microcirculation may result in pulmonary hypertension (PHT) via changes in regional density differences of the pulmonary parenchyma. The aim of this study was to determine whether there were significant differences in CT assessed density gradients in PHT. Fifteen SSc patients, average age 51-6 years (11 women) with depressed gas transfer (TLco < 80% predicted) but with no clinical, functional (forced vital capacity 97% predicted, range 75-125) or high resolution CT evidence of fibrosing alveolitis (FA) were studied. 3/10 mm scans were performed in the prone position. Three circular (20 mm diameter) regions of interest (ROI) were drawn in the most dependent (DL) and non-dependent (NDL) zone of each lung in the upper and lower lobes. Density measures (HU) were selected at random in each ROI. Average DL and NDL densities were calculated in both lobes, and global lung density measurement derived. The density gradient was calculated as the difference in average density measures (DL v NDL) and was assessed in the upper and lower lobes, and as a global lung measure. PHT was defined as an estimated pulmonary artery systolic pressure of greater than 30 mm Hg at Doppler echocardiography. Five of 15 patients had evidence of PHT on echocardiography whilst 10 did not (NPHT). The lung density gradient was significantly smaller in the PHT group than in the non-PHT group. The lower lobes density gradient was 7 (range 4-12) in the PHT group and 12 (2-34) in the non-PHT group (p = 0.05). Similar changes were noted as a global lung measure (p = 0.05). We conclude that in patients with lone PHT complicating SSc there is a significant diminution of the parenchymal density gradient between dependent and non-dependent lung. Determination of CT density gradients may help in the identification of patients with lone PHT. [Supported by a grant from the Raynauds and Scleroderma Association UK.]
A prospective audit of the inpatient management of patients with chronic airflow obstruction

RM ANGS, RK KEITH, JW RAY, RD MONIE, NC THOMSON, KR PATEL  Department of Respiratory Medicine, Western Infirmary/Gartnavel General Hospital and Southern General Hospital, Glasgow In a prospective audit we have examined the care of patients admitted with chronic obstructive pulmonary disease (COPD) to the three major hospitals serving the West of Glasgow. One in two admissions between May 1992 and 1993 were audited, evaluable data being obtained in 261 patients. The main outcome measurements consisted of the use of routine respiratory investigations and standard therapies, the length of stay, inpatient deaths, follow up and readmission rates. Home visits were also performed at two weeks and three months from discharge to assess quality of life using the hospital anxiety and depression (HAD) score and the St George's Respiratory Questionnaire. Lung function was assessed using spirometry. The mean (SD) age of patients admitted was 70.1 (8.8), 89% were smokers with 122 (47%) being male. Arterial blood gases were obtained on admission in 242 (93%) of patients; mean (SD) PaO2, 9.3 (4.6) kPa, PaCO2, 6.3 (2.0) kPa, H+ 41-1 (8-2). One hundred and ten (42%) had a PaO2 of <8 kPa; 97% had a chest radiograph, 70% current or previous spirometry and 47% peak expiratory flow rates measured. Ninety five percent of patients received β, agonists, 83% anticholinergics, 31% theophyllines, 63% corticosteroids, 75% oxygen and 81% antibiotics. Doxapram was used in 7% of patients and 11 patients were given ventilatory support, nine receiving nasal ventilation and two being intubated. The mean (SD) length of stay was 11 (8-5) days with an acute mortality of 17%. Sixty one percent of patients were offered outpatient appointments. Home visit HAD scores indicated significant anxiety and depression, mean (SD) score being 19.8 (7.4) and 18.8 (7.4) at two weeks and three months respectively (NS). Total St George's scores revealed considerable disability with 60 (14%) and 60 (13-1)% respectively (NS). Spirometry was poor, mean (SD) FEV1,0 of 0.77 (0-40) and 0.74 (0-34) at two weeks and three months respectively (NS). By 90 days there were a further 26 deaths giving an overall three month mortality of 27%. Readmission rates were high with 34% being readmitted by 90 days. These findings highlight the need for new strategies to reduce readmission rates and to improve the quality of life of these patients.

Explantation to patients with COPD improves quality of life after discharge

LM OSWAN, A ENGLAND, JAR FRIEND, JG DODD, JA PEARCE  Thoracic Medicine Unit, City Hospital, Aberdeen  Four hundred and seventy-two patients admitted to COPD to chest and general wards were followed up by mail questionnaire within two weeks of discharge. At follow up patients were asked if their hospital admission had been worthwhile (93% agreed) and if their chest was better than before their admission (75% agreed). Patients were also asked if they thought their chest problem had been explained to them while they were in hospital; 74% (225) thought they had had a good or adequate explanation, 13% (39) had an explanation but had not understood it, and 13% (30) had not had an explanation. Older patients were less likely to have had an explanation (p=0.04). Patients who were satisfied with their problem had been explained were more likely to say their chest was better at follow up (79%). Of those with no explanation, 68% thought their chest was better at follow up. With a poor explanation 62% thought their chest was better (p=0.04). A visual analogue scale from 1 to 10 where 1 was “not breathless at all” and 10 was “extremely breathless” was used to measure breathlessness during admission (mean (SE) 5.0 (0-2)) and at follow up (5.7 (0-2), p=0.02). Controlling for age, patients with an adequate explanation had better visual analogue breathlessness scores at follow up than those who had not understood or had not been explained (5.3 ± 5.7) (p=0.05). Good explanation to patients is associated with well being and smaller deterioration in perceived breathlessness after discharge.

Twelve months follow up of autonomic neuropathy in patients with severe COPD

W BIERNACKI, S SCOLHEY, G BOAR, MD PEAKE  Chest Unit, General Infirmary, Pontefract, West Yorkshire  We have shown that subclinical autonomic neuropathy is common in patients with severe COPD (Biernacki et al. Am J Respir Crit Care Med 1994;149-A1012). In one study in diabetic patients deterioration or lack of improvement of autonomic function was associated with an increased mortality rate of observation (Ewing et al. Q J Med 1980;193:95). There are no published data on the natural history of autonomic neuropathy in patients with COPD. We have studied a group of 23 patients (11 M) aged 67 (7) years with severe COPD, FEV1, 31 (9%) predicted, and Pao2 (9) mmHg, 13 (23%) of whom were on LTOT. We measured full standard respiratory function, blood gases and autonomic nerve function (Ewing and Clark) at entry and after one year. In 57% of patients their neuropathy score had deteriorated, in 30% there was no change, and in 13% there was slight improvement. There was no correlation between change of neuropathy score and either change in FEV1, or Pao2, over the year of study. We concluded that autonomic neuropathy deteriorates in the majority of patients with severe COPD, even those receiving LTOT.

Review of the prescription of cylinder oxygen in Scotland

E SKWARSKI, K SKWARSKI, W MACGEE  Respiratory Medicine Unit, Department of Medicine, Royal Infirmary, Edinburgh; Involve Clinical Research, Edinburgh  Long term oxygen therapy (LTOT) can be prescribed in the form of oxygen concentrators or oxygen cylinders. In Scotland concentrators can only be prescribed by chest physicians according to standard guidelines. Cylinder oxygen, which is mainly prescribed by general practitioners (GPs), is cheaper and more convenient. The main reason for this was to determine the characteristics of patients prescribed cylinder oxygen throughout Scotland. From information given by Common Services Agency 791 practices in Scotland were contacted. The response rate was 57% giving information on 746 patients (407 M; mean age 69, range 1-99 years). COPD was the diagnosis in 74% of patients. Oxygen was prescribed for use as required in 82% of patients; on average 8 (range 1-120) cylinders/month were prescribed; 11% of patients used >20 cylinders/month; 27% smoked actively at the time when oxygen was prescribed. Ninety patients (age 69 years, range 36-91 years, 46 M) were visited at home after written consent was obtained from the patient’s GP. In 80% of cases COPD and gradual worsening of dyspnoea was the main reason for which oxygen was prescribed; 40% smoked at the time oxygen was prescribed and 70% had been advised to use oxygen when required. They used a mean of 13 (range 1-120) cylinders/month. Their FEV1 was 0.75 (0-4-1), FVC 1.5 (0-6). A reversibility test (to 5 mg nebulised terbutaline) was performed at home and showed a 15% improvement in FEV1, and an absolute improvement of 200 ml in 16% of patients. SaO2 on air, measured by portable pulse oximeter, was 90 (11%). During daily activities usually causing breathlessness SaO2 fell to 88 (13) (p<0.01). Oxygen consumption (p<0.001) of patients being desaturated by >5% from the baseline during exercise at home. In 1993 £2 million was spent on cylinder oxygen in Scotland in contrast to £350 000 on the oxygen concentrator service. This review highlights the need for guidelines for the prescription of cylinder oxygen. [Supported by Scottish Home and Health Department.]

Early effect of long term oxygen therapy on quality of life in patients with COPD

AA OKUBADEJO, PJ JONES, JA WEDZICHA  Department of Thoracic Medicine, London Chest Hospital, London; Division of Physiological Medicine, St George’s Hospital, London  Long term oxygen therapy (LTOT) has important physiological benefits in patients with COPD and severe hypoxaemia. These patients have impaired quality of life, but it is not known whether there are any early effects of LTOT on life quality. We studied 23 patients (15 F) of median age 71 years with COPD and hypoxaemia, with mean (SD) Pao2, 6-95 (0-75) kPa. PaO2, 6-52 (1-22) kPa, FEV1, 0-75 (0-22). They were assessed on two occasions two weeks apart in the month before starting LTOT, and then a third time two weeks after starting LTOT. We selected a control group of 18 patients (6 F) with COPD and FEV1<1.5 L, but without severe hypoxaemia. Their median age was 72 (range 62-91) years. Patients were offered use of the oxygen concentrator service. This control group had assessments using the same time intervals, but were not started on LTOT. At each assessment both groups underwent spirometry, blood gas analysis, and quality of life assessment using the St George’s Respiratory Questionnaire (SGRQ), and the Hospital Anxiety and Depression Scale (HAD). Before starting LTOT the subjects had a higher mean SGRQ.
total score than the controls (61-9 vs 47-0, p = 0-007), implying a greater impairment in life quality. There were no significant differences between the predicted TLC and FVC assessments in mean score for any mean parameter for either group. At the third assessment after two weeks the mean SGRQ score of the 23 patients fell from 61 to 55-0 (p = 0-045) suggesting improved life quality; however, the control group also experienced a fall (from 45-8 to 41-8) and the mean change was not significantly different between groups (p = 0-8). Changes in anxiety, depression and SIP scores were not significantly different between groups. Although we have demonstrated an improvement in the quality of life of patients after just two weeks of oxygen therapy, this was not significantly greater than that shown by a group of patients without severe hypoxaemia who were not treated with oxygen.

Subjective reactions of patients and their carers to liquid oxygen

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The aim was to investigate the subjective benefits of a portable liquid oxygen (LOX) delivery system for patients and their carers. Patients were included if they satisfied an economic criterion of using two or more cylinders of oxygen per week. Patients were considered suitable if their oxygen saturation fell during a six minute walk and if in the clinician’s view they were motivated to increase activity. One patient declined to use LOX. Of those 44 patients using LOX (22 patients having COPD, 10 having pulmonary fibrosis, and a mixed group of 12 patients with other diagnoses), 33 completed the Oxford Quality of Life (QOL) questionnaire over a period of four months or more. QOL was measured by a specially constructed questionnaire: an activity restriction checklist (including space for other activities to be listed) and a scale measuring the perceived impact of LOX. The activity checklist was administered before and after treatment and the overall evaluation was made monthly after treatment. The mean number of activities restricted prior to LOX was 13-9 (maximum 23, minimum 7), and mean number regained after one month = 5-6 (maximum 17, minimum 1). All patients reported that LOX made their life better, but to different degrees, and all but one carer reported that the carer’s life had also improved. Twenty seven carers felt that LOX had made the patient less dependent, three felt it made no difference, and three felt that the patient had become more dependent on the carer. Patient’s and carers’ evaluations of life improvement were significantly (p < 0.001) correlated (r = 0.64), but patients’ evaluations of life improvement were not significantly different from carers’ evaluations of dependency (r = 0.21). There was a significant improvement in evaluation between the first and second month of LOX use, suggesting that there is a learning component to the advantages gained from LOX.

Factors predicting subjective benefit of patients prescribed liquid oxygen: a pilot study

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The purpose of this study was to investigate whether physiological variables measured prior to the introduction of liquid oxygen (LOX) predicted the degree of subjective benefit after two months treatment with LOX. Thirty one patients were prescribed LOX on the basis of cylinder use (two or more cylinders used per week), oxygen desaturation during the six minute walk test, and the clinician’s opinion that the patient was motivated to increase activity. For all 31 patients FEV1, FVC, TLC, TLoC, KCO, six minute walk distance with oximetry, and resting blood gases were measured pretreatment; the average of patients’ overall evaluation of LOX using a seven-point evaluation scale measured at one and two months after treatment was the outcome variable. Mean (SD) values of physiological variables were: FVC = 0.97 (0.64) l, %predicted FEV1 = 36, FVC = 4.71 (0.81) l, %predicted FVC = 50, mean resting Pco2 = 5.4 (0.96) kPa, mean resting PaO2 = 87 (12-24) kPa, %predicted Kco = 63, %predicted TLC = 84, %predicted TLoC = 43. Pre-exercise oxygen saturation was 93-2%, postexercise oxygen saturation was 86%, mean distance walked was 223.4 m. There was a significant negative correlation between improvement and pretreatment resting Pco2 (r = 0.41, p = 0.03). There were no other significant correlations. Improvement was not significantly related to oxygen desaturation during exercise (r = 0.16). These data suggest that it may be difficult to predict which patients will report most subjective benefit from LOX on the basis of physiological data alone.

Evaluation of the use of concentrators for domiciliary oxygen supply for less than eight hours per day

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Since their introduction in 1985, oxygen concentrators have only been recommended when domiciliary oxygen is used for over eight hours per day. However, concentrators are more cost effective than cylinders when oxygen is used for less than eight hours per day. Twenty six patients in two health districts who used oxygen for less than eight hours per day completed a crossover study in which each group received oxygen from each source for consecutive three month periods. The patients were visited at home before and during the study and on each visit they completed a questionnaire asking about their use of oxygen, how acceptable they found the two sources, and about several dimensions of their quality of life. The theoretical minimum cost of cylinder supply, the actual cost of concentrator supply, and the theoretical concentrator cost were assessed. The patients found the concentrators to be more acceptable, more useful, and less obtrusive than cylinders. They used more oxygen in more of the rooms of the home during treatment with concentrators and there were improvements in the quality of life measurements. The costing information showed that, both in theory and in practice, oxygen concentrators are more cost effective than cylinders when used for less than 1-4 hours per day of oxygen are used. These results suggest that the provision for the supply of domiciliary oxygen should be reviewed and that concentrators should be considered for patients who use more than around 1-4 hours per day.

Use of oxygen concentrators along with nebulisers in the Birmingham and Solihull areas: who are using them and why?

JAYRES, P OTTAWAY Chest Research Institute, Birmingham Heartlands Hospital; Deltibhis Health Care Many patients who use an oxygen concentrator may also be using nebulisers. We set out to determine how many patients were using both nebulisers and oxygen concentrators to assess whether there would be a benefit in trying to develop a combined concentrator and nebuliser. In May 1994 52% of patients in the Birmingham and Solihull areas used oxygen concentrators were sent a questionnaire asking for information on diagnosis, nebuliser use, and other factors concerning the machine and their use. 224 (70%) of questionnaires were returned after a single mailing. Nine patients had died, three forms were not completed, and one patient had moved, leaving 211 for analysis. Eight were being used by children and these are excluded from the following analysis. The mean age (range) of the remaining 203 patients was 65.5 (21–88) years. Eighty four (41%) were women. 148 (73%) had COPD or an equivalent diagnosis and 20 (10%) had a diagnosis of asthma. Thirty (15%) admitted to being a current smoker. One hundred and five (51%) had a nebuliser at home which they used at least once a day. Four of the nebuliser group had a “nondisabling” disease for having a concentrator compared with 28 of 98 in the non-nebuliser group, five of whom gave as their reason “heart failure”. We did not compare as to how many had had formal or informal blood gas assessment. Problems identified by the patients with respect to nebulisers were noise (29 of 105, 28%) and time taken (10 of 105, 9.5%), while 45 of 203 (22.2%) complained of concentrator noise. If these figures can be applied to the rest of the West Midlands, over 500 patients would be candidates for a combined concentrator and nebuliser. Assessment of a prototype combined machine is underway.

Home nebulisers: can optimal treatment be predicted by responses to St George’s Respiratory Questionnaire?

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Domiciliary nebulisers are in widespread use for the treatment of patients with severe chronic asthma or chronic obstructive pulmonary disease (COPD), to relieve symptoms and improve the quality of life. It has been suggested that a 15% improvement in the home peak expiratory flow rate (PEFR) along with a subjective preference for nebuliser treatment after a domiciliary trial can identify patients who should receive long term nebuliser treatment. We tested the hypothesis that this combination of objective and subjective benefits would be reflected in a more formal measure of respiratory related quality of life. We recorded patients’ responses to the St George’s Respiratory Questionnaire (SGRQ) in our nebuliser trials at baseline (Q1), at six weeks (at trial end, Q2) and after 12-16 weeks (Q3). We...
Tumour necrosis factor alpha (TNFα) regulation by interleukin 10 (IL-10) in alveolar macrophages from patients with sarcoidosis (S) or fibrosing alveolitis (CFA)

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We are interested in the regulation of the pro-inflammatory response, in particular the regulation of TNFα by IL-10. Abnormalities of these regulatory mechanisms have been proposed to occur in sarcoidosis (S) and CFA. In a previous study (Am Rev Respir Dis 1994;149:A1066) we demonstrated that IL-10 is a potent inhibitor of TNFα release from human normal alveolar macrophages (AM) and peripheral blood monocytes (PBMM). We hypothesised that in AMs from S or CFA, IL-10 inhibition of TNFα production may be deficient. Bronchoalveolar lavage and peritoneal lavage were performed on all subjects. Recovered AM and PBMM were cultured for a period of 24 hours in the presence of 10 μg/ml lipopolysaccharide (LPS) along with either IL-10 (DNA synthesis, range 0-100 ng/ml) or 10 ng/ml LPS. The free unbound TNFα levels in the AM supernatants were determined by ELISA. There were no significant differences between basal or LPS-induced AM TNFα levels from normals, CFA or S, although the LPS-induced values were lower for S (1.479 (0.2-424)) than for normal subjects (3.508 (0.692), p = 0.071). In normal subjects (n = 6) LPS-induced TNFα levels were significantly reduced in comparison to control values (p<0.01) by a minimum dose of 50 μg/ml IL-10, whereas LPS-induced TNFα levels from sarcoid AM (n = 12) were not significantly affected by IL-10 at any dose. In contrast LPS-induced TNFα levels from CFA (n = 10) were significantly reduced by a minimum dose of 0.1 μg/ml IL-10. IL-10 inhibition of LPS-induced TNFα from PBMM was similar in all three groups. These results suggest an enhanced sensitivity of CFA AM to IL-10, contrasting with the apparent insensitivity of sarcoid AM, either to LPS induction of TNFα or reduction by IL-10. Thus, IL-10 may have a complex role in the regulation of pulmonary TNFα in the inflammatory response.

Hypthesis of growth factor-stimulated tyrosine phosphorylation induces apoptosis in small cell lung cancer cell lines

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Small cell lung cancer (SCLC) constitutes 25% of all lung cancers and follows an aggressive clinical course. Novel approaches to treatment are needed. It is likely that these will arise from a better understanding of the intracellular molecular events underlying mitogenesis. SCLC cells secrete many hormones and neuropeptides. We have previously shown that multiple neuropeptides induce inositol phosphate hydrolysis, mobilisation of intracellular calcium and stimulate growth, suggesting that SCLC growth is mediated by an extensive network of autocrine and paracrine interactions. There is increasing interest in protein tyrosine phosphorylation as a growth promoting signal and we therefore studied the effects of insulin, bombesin and gastrin, all of which are well recognised growth factors, on tyrosine phosphorylation in SCLC. Following stimulation with growth factors, whole cell lysates were analysed by Western blotting. We demonstrated that these growth factors were all able to induce protein tyrosine phosphorylation in SCLC in a concentration and time-dependent fashion. An increase in tyrosine phosphorylation was evident at 30 seconds increasing to a maximum at 10-20 minutes and then declining. Though the pattern of tyrosine phosphorylation was agonist specific, some major substrates like p125FAK and src-related kinases were phosphorylated by all growth factors. Tyrophostin 25, a specific tyrosine kinase inhibitor, inhibits this tyrosine phosphorylation, SCLC growth and stimulates apoptosis in the μM range (IC50 10 μM).

Raised interleukin 10 (IL-10) levels in bronchoalveolar lavage (BAL) samples in the adult respiratory distress syndrome (ARDS)

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ARDS is a catastrophic form of acute inflammatory lung injury. IL-10 plays an important immunomodulatory role in macrophage function. It is a potent inhibitor of the proinflammatory cytokines (e.g IL-1, TNFα) from monocytes/macrophages while at the same time stimulating the release of mediators with anti-inflammation activity (e.g IL-1ra). In this study we investigated whether the level of IL-10 within the alveolar spaces was of prognostic significance. BAL sampling was performed on established ARDS patients within 24 hours of diagnosis. Patient group data are expressed as median values and analysed using Mann-Whitney. IL-10 assays were performed using a sandwich ELISA. Twenty-seven ARDS patients (9/27 died) and a ventilated trauma control group of nine (4/9 died) were enrolled. IL-10 values for patient groups are summarised in the table. BAL IL-10 values were significantly raised
ARDS patients compared with controls (p = 0.0005). In addition, low values correlated significantly with patient mortality (p < 0.01) and a lower PaO2/FiO2 (p < 0.05). Thus, significantly raised levels of the anti-inflammatory cytokine IL-10 were found in BAL samples from ARDS patients. In addition, high IL-10 levels correlated with patient survival. Our data support the hypothesis that an inability to mount a significant intrapulmonary localised anti-inflammatory cytokine response contributes towards a worse overall prognosis.

Plasma thrombomodulin (Tm) in patients “at risk” and with established adult respiratory distress syndrome (ARDS)

PT REID, SC DONNELLY, L BRUCE, AM FERRIE, CR ROBERTSON, TS GRANT, DC CARTER, IR MACGREGOR, C HASLETT Respiratory Medicine Unit, Departments of Surgery, Accident & Emergency, Intensive Therapy Unit; Scottish National Blood Transfusion Service, Edinburgh ARDS represents the pulmonary component of an acute inflammatory microvascular injury in response to a variety of disparate aetiological conditions. Tm is an endothelial surface glycoprotein which may be cleaved and exist in the circulation in a soluble form (sTm). Raised levels have been described in a variety of disease states characterised by damage to the vascular endothelium. In this study we investigated whether significant circulating levels of sTm existed in established ARDS and whether levels in the “at-risk” period had any predictive role for ARDS. Fifty eight “at risk” patients, 16 with established ARDS and a ventilated control group of eight patients were enrolled. Blood sampling was performed on initial hospital presentation for “at risk” and within 24 hours of diagnosis for ARDS patients. sTm was assayed by a sandwich ELISA technique. Non-parametric statistical methods were employed. Significantly higher levels of sTm were found in patients with established ARDS than in controls (p = 0.0001) or those at risk of ARDS (p = 0.001). No significant difference was found in initial sTm levels between patients who progressed to ARDS compared with those who did not (p < 0.1). No significant association was found with subsequent mortality (p = 0.2), ventilatory days (p = 0.06) or organ failure score (p = 0.38). Group patient data are summarised in the table. Our data suggest that sTm provides a marker of endothelial damage in established ARDS but does not provide a marker of impending ARDS in the initial blood sample from patients “at risk” of ARDS.

Lung albumin distribution and content in normobaric and hyperbaric hyperoxia in the rat

PW JOHNSTON, ST NG, AD MCKENNIN, J WOO, JAS BOSS Departments of Pathology and Environmental and Occupational Medicine, University of Aberdeen, Foresterhill, Aberdeen In oxidative lung injury pulmonary oedema results from increased permeability. To assess the severity of injury and its relation to pressure, it would be useful to measure lung leak of a marker. Four groups of rats (n = 8) were exposed to 21 kPa and 100 kPa of oxygen at two or 30 atmospheres absolute (atm abs) with equivalent decompression schedules. Using a new method of image analysis of tissue sections stained with antibody against endogenous albumin, we assessed both the distribution and amount of albumin in resin sections stained with an immunogold method for rat albumin and enhanced with silver for light microscopy. Area fraction (median and interquartile ranges) of positive albumin staining in 100 kPa oxygen exposed groups was greater than 21 kPa oxygen groups regardless of ambient pressure: 1 kPa 2 atm abs, 0.65% (0.60-0.78%); 100 kPa 2 atm abs, 2.85% (2.28-3.53%); 21 kPa 31 atm abs, 0.60% (0.3-1.20%); 100 kPa 31 atm abs, 2.45%, (1.93-3.05%). Albumin, of albumin, corrected for body weight, showed large increases in the hyperoxic groups, again regardless of exposure pressure: 21 kPa 2 atm abs, 53.2 µg/g (42.4-60.2 µg/g); 100 kPa 2 atm abs, 166 µg/g (136-172 µg/g); 21 kPa 31 atm abs, 48.0 µg/g (3.2-87.7 µg/g); 100 kPa 31 atm abs, 151 µg/g (126.5-155.0 µg/g). Statistical analysis confirms the differences relate to partial pressure of oxygen and not exposure pressure. We suggest that risks of oxidative injury in hyperoxia result from the partial pressure of oxygen and are not pressure dependent.

4-Hydroxy-2-nonenal levels in the plasma of patients with ARDS

GJ QUINLAN, TW EVANS, JMC GUTTERIDGE Royal Brompton Hospital, London Recent findings have shown that the n-6 fatty acid oxidation product, 4-hydroxy-2-nonenal (HNE) is increased in the plasma of ARDS patients compared with healthy controls and that such increases are related to losses in the polyunsaturated fatty acid linoleic acid. We have now developed a more sensitive assay of HNE based on gas chromatography mass spectrometry of an HNE trimethylsilyl (TMS) derivative. Results are in agreement with those obtained previously. ARDS non-survivors had higher HNE levels (0.552 (0.069 nmol/ml) than survivors (0.433 (0.048 nmol/ml), although not significantly (p = 0.6). Comparison of both groups with healthy controls (0.205 (0.03 nmol/ml) showed highly significant (p < 0.0001) differences. In addition, a control group of cardiopulmonary bypass patients was also included (0.279 (0.027 nmol/ml). These were significantly different from the normal control group (p < 0.05) and the ARDS non-survivors (p < 0.03), but not from survivors (p > 0.1). HNE is a lipid peroxidation product with several well documented biological effects such as cytotoxicity, enzyme inhibition and chemotactic properties. Our findings show that ARDS patients are under considerable oxidative stress which results in molecular damage and the formation of toxic oxidation products including HNE. These products may influence the course of ARDS in non-surviving patients. The increased levels of HNE in patient controls compared with healthy controls is probably a reflection of their oxidative stress from ventilatory support. This new assay technique on a larger patient population is in agreement with our previous findings.

Plasma 4-hydroxy-2-nonenal levels during cardiopulmonary bypass and their relationship to the iron loading of transferrin

GJ QUINLAN, S MUMFY, J PEPPER, TW EVANS, JMC GUTTERIDGE Royal Brompton Hospital, London 4-hydroxy-2-nonenal (HNE) is an aldehyde lipid peroxidation product formed in vitro in model systems from the oxidation of n-6 fatty acids. It has also been detected and measured in vivo in plasma and tissue. HNE is a highly reactive molecule which can exert profound influences on biological systems including enzyme inhibition and chemotaxis. Reactive forms of iron have been shown to be essential catalysts for HNE formation, and plasma linoleic acid has been implicated as a substrate for its formation in patients with ARDS. We have measured HNE levels using chromatography mass spectrometry, in 12 cardiopulmonary bypass (CPB) patients before and after bypass. When corrected for haemodilution, by relating results to total protein levels, all bypass patients showed increases in HNE after bypass (11.5 (2.7 pmol/mg protein) compared with levels before bypass (8.5 (2.4) pmol/mg protein). Further, when plasma catalytic iron levels were measured as boro- myeloblastable iron, findings of the 12 patients were found to be iron overloaded after bypass, and it was these patients who showed the greatest percentage increase in HNE (90.6% (16.85)) compared with patients not iron overloaded (21.6% (5.98), differences (p = 0.01). These results indicate that CPB patients are under oxidant stress as a consequence of extracorporeal (p < 0.03) and iron, which can release catalytic iron and exacerbate oxidative damage. CPB patients are also subjected to further oxidative stress from ischaemia/reperfusion injury associated with cross clamp release, and reactive iron will increase oxidative damage. Our results support the use of chelation therapy during bypass.

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Temporal pattern of enhanced expression of procollagen-1 in tuberculin induced delayed type hypersensitivity (DTH)

BG Marshall, A Wangoo, HT Cook, RJ Shaw Departments of Respiratory Medicine and Histopathology, St Mary's Hospital Medical School, London Tissue fibrosis is a common and serious consequence of chronic inflammation in lungs of patients with tuberculosis. Mechanisms linking these processes are poorly understood. In this study we have used the human in vivo model of the tuberculin Heaf test to define the temporal sequence of events leading to the increase in new collagen production, since this would provide important evidence that a DTH response can progress to a fibrotic reaction. Thirty-six volunteers with grade 3/4 Heaf reactions (six groups of six) underwent a skin biopsy at the site of tuberculin at intervals from day 1 to 14. A group of 12 controls received normal saline. Immunohistochemical analysis revealed increased expression of procollagen peptide 1 (PCP-1) as early as day 3 following the tuberculin Heaf test and was maximal on day 14. There was marked expression of PCP-1 in the perivascular infiltrates and in the surrounding interstitial infiltrate. These data from a human in vivo model of a delayed type hypersensitivity response provide information on the course of events in this type of chronic inflammation, and indicate that the immune response is intimately associated with the initiation of a fibrotic response.

Tuberculosis among the homeless in London

D Kumar, RK Citron, JL Leese, J Watson PHLS Communicable Disease Surveillance Centre (CDSC); Royal Brompton Hospital; Department of Health Tuberculosis is a well recognised problem among the homeless. Recent figures show that tuberculosis notifications are no longer declining in England and Wales, with an increase of around 5% per year in the last two years. An increase in homelessness has been suggested as one of the possible contributors to this increase. Reliable estimates of the prevalence or incidence of tuberculosis in the homeless subgroup are not available for this country. A study was undertaken to estimate the prevalence of active pulmonary tuberculosis in a homeless population in London and to assess whether those with suspected disease could be integrated into the existing health care system for further follow up and treatment. Voluntary screening programmes based on a questionnaire survey and chest radiography were set up during Christmas 1992 and 1993 at temporary shelters for the homeless in London. In 1992 nearly 1600 people visited the centre, of whom 372 volunteered for the screening and 342 were x rayed. Nineteen (5.4%) of those x rayed had radiological features suggestive of active tuberculosis. In 1993 around 2000 homeless people visited the centre, of whom 270 volunteered for the screening and 253 were x rayed. Eleven (4.4%) had features consistent with active tuberculosis on the basis of the chest x rays and clinical examination by a chest physician. Overall, among 595 people x rayed in the two surveys 30 (5%) had changes suggestive of active tuberculosis. Further investigations confirmed nine (1.5%) with active pulmonary disease and eight with no active tuberculosis. In 13 the diagnosis was not determined as four declined further investigation and nine did not attend their hospital appointment.

Trends in tuberculosis notifications in England and Wales, 1982–93

AG Hayward, JM Watson PHLS Communicable Disease Surveillance Centre, London Between 1987 and 1993 over 10 000 more cases of tuberculosis (TB) were notified than would have been expected from previous trends had continued. TB notifications from 1982 to 1993 were analysed to determine the population groups and geographical areas in which increases occurred. Three year average notification rates for the years 1986–8 were compared with those for 1991–3 (or 1990–2 when notification data were unavailable for 1993) in order to calculate comparative rates (CR within 95% confidence limits). Increases were greatest in Regional Health Authorities (RHAs) with large urban concentrations (North East Thames: CR = 3.33 (1.25–4.1)); West Midlands (CR = 1.12 (1.05–1.18)). Rates in rural RHAs generally continued to decline (Wales CR = 0.92 (0.91–0.93); Wessex CR = 0.81 (0.70–0.93)). In men significant increases were seen in those aged 15–25 years (CR = 1.2 (1.0–1.32) and decreases in those over 35 years. In women no age groups experienced significant changes in rates. Non-respiratory TB accounted for 21% of notifications in 1987 and 27% in 1993. In most age/sex/geographical groups significant increases were confined to non-respiratory TB. However, increases were seen in both respiratory and non-respiratory rates (CR = 1.24 and 1.28 respectively) in men aged 15–34 years in North Thames RHA. Some of the increases observed may be due to increased reporting of TB cases following the appointment of Consultants in Communicable Disease Control. This is supported by selective increases in non-respiratory notifications which may have been subject to greater under notification (than respiratory TB) in the past. Other factors associated with recent increases will be discussed. In addition the results of the 1993 National Tuberculosis Survey, which includes information on ethnicity, will further elucidate the causes of the increases.

More ambiguities and inaccuracies in the notification of cases of tuberculosis in England and Wales

D Kumar, J Watson, J Darbyshire on behalf of a PHLS/BTH collaborative group PHLS Communicable Disease Surveillance Centre, London; MRC HIV Clinical Trials Centre, Royal Brompton Hospital, London There has been a statutory requirement to notify all cases of tuberculosis to the clinical suspicion, since 1973. "Ambiguities and inaccuracies" in the notification of tuberculosis were identified in the 1978/79 MRC TuberculosisNotifications Survey and addressed in a Code of Practice for notification published by the British Thoracic Association in 1982. The 1993 National Tuberculosis Survey has revealed that problems in the reporting of tuberculosis, both old and new, remain. Copies of 6118 notification forms were received in the 1993 National Tuberculosis Survey of which 550 were duplicates (117 cases were reported by two different local authorities and 140 were reported twice by the same local authority). Five hundred and sixty three cases receiving chemoprophylaxis only were notified; in 425 this was stated on the notification form (and could therefore be excluded from statistics prepared by the Office of Population Censuses and Surveys), but in the remaining 138 chemoprophylaxis was only apparent from the survey clinical form. In 280 cases the diagnosis was changed after notification, but only in around 50% of these is there evidence that the case was denotified. In 145 cases (mainly children) it was reported that two drugs only were to be given (for longer than three months) and it was uncertain if
Tuberculosis incidence in Scotland is increasing in the young and the elderly

JS DUFFIELD, WH ADAMS, M ANDERSON, AG LEITCH  Respiratory Medicine Unit, Royal Victoria Chest Clinic, RIE NHS Trust; Medical Statistics Unit, University of Edinburgh, Edinburgh  Notifications of tuberculosis by age, sex and site of disease for the years 1981–92 were obtained from the Information and Statistics Division of the NHS in Scotland. Total, as well as age and sex related incidence rates, were calculated for each year using General Register Office population estimates. Total tuberculosis incidence rates per 10^5 population for 1981–92 are shown in the table and demonstrate a decline succeeded by a plateau in 1987. Age group analysis of pulmonary tuberculosis showed that incidence (10^−5) decreased from 7-4 in 1981 to 2-6 in 1987, rising significantly (p=0.012) by an estimated 12-6% per annum to 3-7 in the 1990–92 age group. In the 65+ age group incidence declined from 30-1 in 1981 to 17-3 in 1988 and rose significantly (p=0.048) by estimated 4-1% per annum to 22-2 in 1992. Age groups 15–44 and 45–64 show a continuous decrease in rate on natural logarithmic scales from 11-2 and 23-8 in 1981 to 4-3 and 11-4 respectively in 1992. Inclusion of extrapulmonary cases to give total tuberculosis notification rates revealed almost identical findings within all age groups with the same patterns on regression analysis of incidence. The estimated annual increase in total incidence in 1992 was 8-5% (p=0.035) for the 0–14 age group and 2-8% (p=0.0102) for the 65+ age group. Pulmonary tuberculosis in Scotland is increasing significantly in the young and the elderly; the increases may be causally related.

Characteristics and outcome of patients with pulmonary non-tuberculous mycobacterial infections

L DAVIES, PDO DAVIES  Tuberculosis Research Unit, Aintree Chest Centre, Fazakerley Hospital, Liverpool  Studies of patients with non-tuberculous mycobacterial infections have shown a high mortality. We performed a retrospective case study of all patients presenting to Aintree Hospitals, Liverpool over a six year period from January 1988 to December 1993 with culture proven non-tuberculous mycobacterial disease. There were 38 patients in total; seven were HIV positive and four had non-respiratory infections. These were excluded. Of the 27 remaining patients 18 (67%) were men (age range 44–84, mean 66 years). A total of 19 subjects smoked at the time of diagnosis, seven were ex-smokers and one only was a lifelong non-smoker. Seven drank more than twice the recommended limit of units of alcohol per week. Fifteen patients had COPD with FEV1, values ranging from 24–84% of predicted (mean 55%). Twelve (44%) had evidence of other smoking related disease and nine (33%) had a history of M tuberculosis disease. At least two isolates were obtained from each patient, in 26 cases from sputum alone and, in the remaining patient, from bronchial washings. Fifteen (56%) grew M kansasii, six (22%) MAC, five (18%) M maltosone, and one (4%) M xenopi. Of the 19 who smoked at the time of diagnosis, seven (37%) died within the study period (four reportedly of COPD), three remained unwell with frequent admissions due to cough and breathlessness, six are now well, and three were lost to follow up. Of the seven who were ex-smokers at the time of diagnosis four (57%) died within the study period (three reportedly of COPD and one of M kansasii), one remains breathless, one is now well, and one is lost to follow up. The one lifelong non-smoker responded rapidly to treatment and is now well.

Non-tuberculous mycobacterial infection appears to be predominantly a smoker’s disease.

Tuberculosis screening and tuberculin testing in two social clubs in the city of Liverpool

PDO DAVIES, CSD WILLIAMS, S JAMESON, A STEEL, Q SYED, M BELLIS, N BEECHING  Tuberculosis Research Unit, Regional Epidemiological Centre, Infectious Diseases Unit, Aintree Hospitals, Liverpool  In the course of an investigation into an outbreak of tuberculosis (TB) in a social club in Liverpool in 1993 198 club members were screened by chest radiography and Heaf testing. All those with grade 3 and 4 were offered a chest radiograph. No further cases of TB were found. In order to determine the background population of Heaf positivity within Liverpool a controlled study was done in a social club in a part of Liverpool where no cases of TB had been reported. 171 "controls" were tested. Percentage of Heaf grade for the two clubs is shown in the table. The control population was older than those from the TB affected club. There was no statistical difference between those that had received BCG vaccination and those who had not. The club where TB cases had been reported showed a significantly greater number of members who were Heaf grades 3 or 4 (<p<0.005). The study at the "control" club showed that a significant proportion (27%) of adults are likely to have had previous TB infection. Adults attending the social club where there had been TB cases reported showed a higher proportion (41%) of members with Heaf grades 3 and 4. It is probable that the difference in proportion of grade 3 and 4 test positive subjects (14%) represents recent infection.

Tuberculin test screening in substance abusers

PDO DAVIES, C MORRISON, D WAKEFIELD, SM RUBEN, N BEECHING  Tuberculosis Research Unit and Maryland Centre, Liverpool  Drug users have been one of the most at risk groups from HIV infection due to their lifestyle through either the sharing of contaminated injecting equipment or sexual behaviour. Tuberculosis (TB) is often associated. Reports from the USA recommend that TB screening should be undertaken in substance abuse treatment centres as drug users will be at high risk because of the higher seroprevalence of HIV and adverse social factors. The Liverpool Drug Dependency Clinic is the largest clinic in Europe, treating over 1000 opiate addicts each year on a methadone programme. One hundred and eighteen drug users had Heaf testing performed and read within 10 days. The mean (SD) age was 28-4 (5-6) years (range 17–46). There were 77 males (65% (91)) and 41 females (35% (118)). Ten (8-4%) were homeless and 21 (17-8%) lived in households of five or more. Eighteen (15-3%) reported having a family member who had had TB. There were four non-smokers and 70 non-drinkers, 17 (14-4%) drinking over 35 units a week. Seventeen women had exchanged sex for money. In the previous four weeks 52 (44%) had injected an illicit drug. Nineteen (16%) had smoked Crack. Heaf test grade results showed that 26 (22%) were grade 0, 85 (72%) grade 1 or 2, and seven (6-0%) grade 3 or 4. Sex, age, homelessness, employment status, smoking, alcohol or drug use were not significantly associated with any Heaf grade result. Forty three (36-4%) had never had school BCG vaccination and 20 of these had a grade 0 result. Only previous BCG was associated with a positive Heaf test (<p<0.001). Only five of the 33 referred for BCG vaccination or further assessment at the nightclub clinic attended. Risk of TB infection amongst Liverpool substance abusers is low.

Experience of miliary tuberculosis in a high incidence area: Blackburn 1978–93

LP ORMEROD  Chest Clinic, Blackburn Royal Infirmary, Blackburn Lancs  Thirty nine cases of miliary tuberculosis (25 cryptic: 14...
classical) were seen in a population of 265,000 with over 10% of Indian subcontinent (ISC) ethnic origin over a 16 year period, giving an annual incidence of 0.92/100,000. Of the 14 classical cases there were white (mean age 51.3 years; range 2–79), three Pakistani (mean age 55.7; range 31–53), and one each from India (mean age 56) and Indonesia (mean age 36). Five (36%) had associated TB meningitis with positive CSF cultures, five had positive sputum cultures and one positive histology from a skin lesion. Two died on treatment at four days (age 93, TBM) and six weeks (age 23), after commencement of therapy. Of the cryptic cases seven were white (mean age 53.7; range 12–76), eight ISC (mean age 39.5; range 16–85) and eight Indian (mean age 39.5; range 21–71). Twenty two of 25 (88%) had a normal chest radiograph, one had old apical calcification, and two focal nodular pulmonary TB. Three patients (white aged 48, Pakistani aged 38, Indian aged 31) were all diagnosed pneumothorax examination following admission to the outpatient clinic. One other case was confirmed by positive sputum culture, the rest showed response to specific antituberculosis treatment and fulfilled Proudfoot’s criteria (Proudfoot et al. BMJ 1969;i:273–6).

The outcome with 5/39 (13%) mortality and 25/39 (64%) cryptic miliary TB is substantially different from the spectrum and outcome of disease reported from Edinburgh (Simé et al. Am Rev Respir Dis 1993;147:A121) which has a minimal immigrant community. The better outcome in a high prevalence, high immigrant age group probably reflects greater awareness of the diagnosis and prompter initiation of treatment.

Effects of winter pollution on respiratory symptoms and medication use in subjects with asthma and chronic bronchitis

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We have previously reported on the effects of wintertime air pollution on measurements of peak flow in subjects with asthma and chronic bronchitis (Thorax 1994;49:360P). Here we report on the results of the analysis of symptoms and medication use in the same group of subjects. For one month during January–February 1992 subjects recorded their daily bronchodilator use (BU) and symptoms of wheeze, dyspnoea, cough, throat and eye irritation were scored on a visual analogue scale. Daily means of SO2, NO2, and O3 were measured by absorption spectroscopy. After allowing for between subject differences and controlling for temperature, multiple regression analysis was used to investigate whether subjects complained of wheeze, etc and used more than their average BU as levels of same day, 24 and 48 hour lags of pollution increased. Satisfactory records were received from 30 subjects (21 reactors with methacholine PD10, FEV1 <125 µmol identified). Max daily mean SO2 and NO2 (169, 151 µg/m3) exceeded guidelines for health, O3 41 µg/m3. Same day O3 was associated with increased BU in all subjects (p<0.01) and in reactors a trend was observed between 48 hour lag O3 and dyspnoea (p<0.06). No other adverse effects were found. We conclude that, at the levels encountered, the winter pollution must have little effect on respiratory symptoms in this group of potentially more sensitive individuals. As in a previous summer study (Am Rev Respir Dis 1993;147:A637) BU is again associated with ambient O3 levels.

Toxicity of environmental particulate: studies with ultrafine titanium dioxide

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Environmental studies have demonstrated a firm relationship between the levels of fine particles suspended in environmental air (PM10) and (a) respiratory and cardiovascular mortality; (b) attacks of asthma and COPD. PM10 particulate cannot be obtained in sufficient quantities for experimentation and so we utilized two samples of the "inert" pigment titanium dioxide: normal TiO2 (NTiO2) 250 nm diameter and ultrafine TiO2 (UFTiO2) 25 nm diameter (Ferin et al. Am J Respir Cell Mol Biol 1992;6:535–42), the latter being a surrogate for the fine environmental particles. We added NTiO2 and UFTiO2 to cells of an epithelial cell line A549 for four hours and showed that UFTiO2 caused dose-dependent detachment of the cells from their substratum, without toxicity, whilst NTiO2 had very little effect: data as mean (SE) cpm in detached cells to triplicate wells minus control value in four separate experiments: 50 µg/ml NTiO2 82 (144); 50 µg/ml UFTiO2 1018 (607); p<0.05. The ability to nick supercoiled DNA, detected by quantitative scanning laser densitometry of ethidium bromide-stained agarose electrophoresis gels, is a sensitive assay for the ability of particle surfaces to release free radicals. UFTiO2, caused 100% supercoiled DNA strand breakage (three separate experiments) whilst NTiO2 had no effect. The effect of UFTiO2 was greater than we have seen with short chain oxides (50 nm) and hydroxyl radicals. A substantial fraction of environmental particulate is in the ultrafine (<50 nm) size range and, if these have the same activities as shown here for the UFTiO2 then this could help explain why PM10 is so toxic.

Occupational asthma due to a widely used soft solder flux not containing colophony

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Telux flux is a safe alternative to colophony soldering flux, containing zinc and ammonium chlorides as its main active constituents. We describe flux fume induced asthma in a previously well 19 year old female gas appliance servicing engineer. The initial presentation was the development of a viral-like illness after 10 weeks regular work exposure to Telux flux. She then noted several episodes of wheeze and exertional dyspnoea which appeared work related. Her subsequent work environment was changed so as to minimise exposure before further investigation. Atopy was diagnosed (sensitisation to house dust, dust mites, grasses, pets) and in the light of the subject's symptoms a diagnosis of occupational asthma and hayfever were noted. An initial methacholine test (Newcastle protocol) showed a high degree of airway responsiveness (AR) with a PD20 of 10 µg, despite her remaining relatively asymptomatic. Four months after exposure ceased PD20 had markedly increased to 1159 µg. Laboratory based inhalation provocation tests were performed with 30 and then 60 minutes of fume exposure group while soldering. The 30 minute exposure test had the baseline FEV1, falling by 25% but only just meeting the 95% lower confidence interval. The FEV1, area decrement (AD) was increased significantly from 0-06 to 4-92 litre hours (p<0.01) and PD20 fell from 800 to 100 µg. The 60 minute test saw the baseline FEV1, fall by 58% during the late asthmatic reaction (LAR) but again there were minimal symptoms. The area decrement was increased significantly from 0-033 to 21-5 litre hours (p<0.001) and again PD20 fell dramatically from 1706 to 34 µg. Two weeks after the last challenge test PD20 had risen to 2094 µg, a value which is not usually associated with active asthma. Zinc chloride has previously been implicated in soft solder asthma (Thorax 1989;44:220–3) and it is likely that any mineral oils present will burn off in the soldering process. The immediate and late falls in FEV1, and increases in AR demonstrate, despite minimal symptoms, that zinc is capable of inducing occupational asthma. This case demonstrates the wide variation in AR typical of early stages of the illness. We suspect the zinc chloride component of the flux acting as a sensitisation agent in certain individuals.

Influence of a variety of personal characteristics upon across shift and across week changes in lung function in textile workers

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One hundred and ninety six textile workers performed spirometry and bronchial reactivity (BR) before and after their first and fourth working shifts of the week. The effect of several factors upon across shift and across week changes (day 1 pre-shift minus day 4 pre-shift) in spirometry and BR was studied. No differences were found between men and women, whites and Asian subpopulations. Cotton workers (as opposed to manmade-fibre workers (MMF)) had significantly lower percentage of predicted spirometry and significantly higher baseline BR (p<0.05). No differences between the two groups for across shift and across week changes could be demonstrated. A high cotton exposure group demonstrated a significant increase in BR across shift on day 1 but not day 4 (p<0.05), and there were no other differences. Operatives who had ever smoked (ES) showed a significantly lower % predicted spirometry and higher baseline BR than the lifelong non-smoker (NS) group (p<0.05). The ES group demonstrated significantly greater across week falls in all spirometric parameters compared to the NS group. None of the personal characteristics seemed important in the production of across shift changes in lung function; however, cigarette smoking was the most important one involved in the production of changes in spirometry across the working week in these textile workers.
Acclerated lung function decline in textile workers: cotton dust, cigarette smoking or both? C J WARBURTON, AM FLETCHER, CAC PICKERING, RM NIVEN, LA OLDHAM, HC FRANCES 
Northwest Lung Centre, Wythenshawe Hospital, Manchester. One hundred and ninety six Lancashire textile workers performed spirometry before the first shift of the working week as part of an ongoing study of across week variations in lung function. Percentage of predicted lung function was calculated using the reference equations of Quanjer (Bull Eur Physiopathol Respir, 1983) for whites, and our own previously presented regression equations for Asians (Warburton et al. Thorax 1994). Correlations between various predictor parameters and percentage predicted lung function were calculated (SPSS) with 194 degrees of freedom. Significant negative correlations between PPFVEF, and PPFPEF<sub>2.5</sub> and (1) the cumulative number of cigarettes smoked (cig/week), (2) the number of years spent working in the textile industry, and (3) the number of work-related respiratory symptoms present in each operator were found. PPFVEF, PPFVFC and PPFPEF<sub>2.5</sub> were all significantly negatively correlated with age (the influence of which should have been removed by the regression equations). These results suggest that accelerated decline in the spirometric indicators of bronchial obstruction (PFVEF, PPFPEF<sub>2.5</sub>) are related to cigarette smoking and to the number of years of exposure to cotton dust. These two factors may be interrelated and the relative importance of each is difficult to ascertain. The strong correlations with age may suggest that, as operators age, the effects of longer cotton dust exposure and cumulatively more cigarettes are at least additive.

Airborne house dust mite allergen
A CUSTOVIC, SCO TAGGART, A WOODCOCK North West Lung Centre, Wythenshawe Hospital, Manchester. Airborne sampling is widely used in the assessment of occupational exposure and might be considered as a more representative measure of inhaled allergen than reservoir levels. However, its use as a method for house dust mite allergen measurement is still controversial. Personal airborne samples were collected with Casella sampler (21/min): overnight in bed with a modified face mask (8 hours; n = 7); during daily activities (8 hours; n = 7); and during laboratory sieving of domestic dust samples (2.5 hours; n = 8). Fixed location samples were collected with a high volume sampler (60 l/min) before and after disturbance (4 hours; n = 8); overnight with Casella sampler attached to pillow (8 hours; n = 30) [mattress and bedding Der p I: geometric mean (GM) 9320 ng/g. Der p I was undetectable (<1 ng/ml) in all personal airborne samples collected overnight and during the usual domestic activities (mean sample volumes (MSV) 0-93 m<sup>3</sup> and 0-96 m<sup>3</sup>, respectively). High levels of airborne Der p I were measured during sieving of dust (GM 38-9 ng/m<sup>3</sup>, range 18-3-72-6; MSV 0-3 m<sup>3</sup>). Der p I was undetectable in all fixed location samples collected before disturbance (MSV 15-5 m<sup>3</sup>). Detectable levels (>1 ng/ml but <4 ng/ml) in 3/30 and undetectable in 27/30 overnight samples (MSV 0-95 m<sup>3</sup>). As yet, airborne sampling is insufficiently sensitive to produce reliable and repeatable results. More sensitive assays or better sampling methods are needed.

Levels of exposure to mite allergen in hospital wards
AM FLETCHER, A CUSTOVIC, CAC PICKERING, H FRANCES, AA WOODCOCK North West Lung Centre, Wythenshawe Hospital, Manchester. House dust mites are generally accepted as a major source of indoor allergen. In 1994 Mohindra et al. (Clin Exp Allergy 1994;24:174) reported levels of house dust mite allergen recovered from carpeted wards in excess of those considered to be a risk factor for acute asthmatic attacks. In this study three hospitals were investigated. The investigation included four carpeted wards in two NHS hospitals and the carpeted single rooms of a local private hospital. The age of the carpets in the wards varied between 18 months and 10 years. Fine dust samples were collected by vacuuming (flow 45 l/s) 1 m<sup>2</sup> of carpet and mattress for two minutes. The Der p I allergen content was assayed by a two-site immunoenzyme assay. In total 21 mattresses and 26 carpet dust samples were analysed. The mite allergen levels were found to be low in all three hospitals in all sample areas. The geometric mean level of Der p I antigen collected from the mattresses was 234 ng/g (range 100-1200 ng/g) and from the carpets 138 ng/g (range 100-410 ng/g). In one hospital dust was also collected from the surface of six fabric covered chairs situated in the outpatient area. These samples provided the highest levels of mite allergen; geometric mean 975 ng/g (range 680-1200 ng/g). This study demonstrates that mite allergen levels in carpeted hospital wards and on sealed hospital mattresses are at a low level and are unlikely to be of clinical significance to house dust mite sensitive asthmatic patients.

Levels of contamination of domestic dust with endotoxin and Der p I
JCG SIMPSON, A CUSTOVIC, CAC PICKERING, AM FLETCHER, R MCL NIVEN, LA OLDHAM, AA WOODCOCK North West Lung Centre, Wythenshawe Hospital, Manchester. Environmental factors are clearly important in the pathogenesis of asthma. While there is a variety of allergens at home, importance, a wide variety of indoor and outdoor pollutants may have a role in modifying the asthmatic response. Endotoxins are derived from the cell wall of Gram negative bacteria and are potent pro-inflammatory agents. They are found to be contaminants of various occupational environments and are implicated as aetiological agents in a number of diseases (including byssinosis, organic dust toxic syndrome, and humidifier fever). We investigated the contamination of domestic dusts with endotoxin and mite allergen Der p I in 13 houses. Paired fine dust samples were collected by vacuuming (flow 45 l/s) 1 m<sup>2</sup> of mattress and living room carpet for two minutes. Samples were assayed for endotoxin (turbidimetric method using the LAL 5000e machine) and Der p I (two site immunometric ELISA). The mean (SD) level of endotoxin collected from the carpets (expressed as ng/g of dust) was 14.8 (23.9) and was significantly greater than from the mattresses (2.8 (6.1); p = 0.001. Der p I levels (expressed as ng/g of dust) were; carpets 7395 (17 651) and mattresses 5493 (10 195). No correlation between measured endotoxin and Der p I levels was found. This study documents the contamination of the domestic environment with endotoxin. Further work is indicated to ascertain the role of this potential exposure in asthma and specifically to determine any synergistic activity with allergen exposure.

Allergic bronchopulmonary aspergillosis after lung transplantation?
JJ EGAN, N YONAN, KB CARROLL, AK DEIRANIYA, AK WEBB, AA WOODCOCK North West Lung Centre, Wythenshawe Hospital, Manchester. Experience in lung transplantation suggests that asthma is an organ specific condition. We describe two cases of allergic bronchopulmonary aspergillosis following lung transplantation for cystic fibrosis (CF). Patient A: a 37 year old male CF patient received a heart/lung transplant. The 18 year old donor had no history of asthma. Preoperatively the recipient was negative for Aspergillus precipitins (Asp ppts) and he was Aspergillus fumigatus (Asp f) spurt negative. One year after surgery he was weaned off steroids. Three years postoperatively he presented with a wheeze, a peripheral eosinophilia of 4 x 10<sup>9</sup> (normal<4 x 10<sup>9</sup>), skin tests were positive to Asp f only, total IgE = 70 IU/l (normal<200), RAST to Asp f was positive at four units (normal<0.35). HRCT showed areas of consolidation and bronchoscopic mucus plugs were aspirated and Asp f was cultured. He responded to prednisolone and inhaled fluticasone. Patient B: A 20 year old male patient received a double sequential lung transplant for CF. The recipient was atopic with positive skin tests to Der p I, grass and Asp f. Prior to transplant Asp f was not isolated from his sputum and he was Asp ppts positive. The 15 year old donor had mild asthma. Four months after surgery he presented with complete collapse of the left lung. Bronchoscopic recollection mucus plugging of the left bronchial tree. The plugs contained eosinophils and fungal hyphae. He had a peripheral eosinophilia of 1.1 x 10<sup>9</sup> and a total IgE = 1500 IU/l (<200) and RAST to Asp f was 55 units (<35). Methylprednisolone 500 mg for three days and nebulised budesonide daily resulted in a rapid resolution of the radiological and bronchoscopic findings. The patient was discharged from hospital two days covered by oral iraconazole and lypoosomal amphotericin. These cases indicate that both systemic and local mechanisms can contribute to an allergic reaction in the airways.

Audit of an admission data sheet for patients with acute asthma
P HUGHES, C SHEPHERD, CR MCGAVIN The Chest Clinic, Freedom Fields Hospital, Plymouth. Assessment of acute asthma relies on measure-
Audit of emergency self-admission for asthma

VA VORA, JE SCULLION, MDL MORGAN, AJ WARDALE Department of Respiratory Medicine, Glenfield Hospital, Leicester The BTS guidelines for asthma stress the need for an emergency self-admission service for patients at risk. A dedicated emergency bed for this purpose has been provided in Leicester since 1984. To assess the quality and appropriateness of this service we have audited admissions for a five month period between April and August 1993. The admitting nursing and medical staff collected the following data for each admission: interval between notification, arrival and initiation of therapy; mode of transport; reason for admission (related/unrelated to asthma); and assessment of severity by estimation of peak flow (PEF) and arterial blood gases (ABG). Of the 147 patients on the self-admission list, 47 were admitted during the 153 days of the study period. Thus, 106 bed days (70%) remained unoccupied. More than 50% (26/47) transported themselves to hospital. Mean interval between notification and arrival was 52 minutes (R = 5 min to 2-5 hours), and between arrival and initiation of treatment was 20 minutes (R = 0-2-3 hours). Seven patients were admitted with problems unrelated to acute asthma. Arterial blood gas tensions were normal in 17 (36%) patients and PEF was within expected range in two patients. There were no admissions to ITU or fatalities amongst the patients on the self-admission list during the study period. In conclusion, even though self-management plans were not universally applied, 74% of admissions were deemed appropriate by the admitting staff, but the bed occupancy was maintained at 30% (30/100). Therefore the self-admission system was felt to fulfill a need but the cost should be recognised in the contracting process.

New asthma referrals to hospital clinics

I BYLAND, MG PEARSON ON BEHALF OF THE NATIONAL ASTHMA TASK FORCE AND 30 PARTICIPATING PHYSICIANS Aintree Chest Centre, Liverpool There are few data about the patterns of referral to, or what happens in, hospital outpatient clinics. The National Asthma Task Force has surveyed 21 adult and nine paediatric hospitals collecting data on 335 new adult patients and 152 new paediatric referrals. Median age of adults was 43 (13-85) years and children 3 (months-14) years. 57% of adults and 39% of children were female; 259/296 (88%) adults and 121/144 (84%) children were directly referred by the GP. For adults, of the 259 GP letters, 73 (28%) requested help with diagnosis, 132 (51%) asked for advice on treatment, and 22 (8%) indicated patient pressure for referral. In children corresponding figures are 55/122 (45%), 55/122 (45%) and 7/122 (6%), respectively. Treatment on referral together with the treatment recommended after consultation is shown in the table. For adults the first outpatient consultation increased treatment in 106/299 (46%), left it unchanged in 104 (45%), and reduced it in 19 (8%). For children treatment was stepped up in 40%, unchanged in 48%, and down in 12%. Although 52% of adults and 40% of children reported waking with asthma for two or more nights in the last week, the changes in treatment at the first visit by GPs were not correlated with these symptoms. Adult and paediatricians discharged 21% at visit 1 and planned early discharge in another 31%. Most asthma patients are referred by GPs; half have their treatment increased at first visit and early discharge back to GP is planned in 70% of adults and 52% of children. Most physicians intend long term follow up only for the most severe asthma patients. [Supported by a grant from the Department of Health.]

Asthma and poverty in the West Midlands

JP WATSON, P COWAN, RA LEWIS Worcester Royal Infirmary, Worcester The suspected link between asthma and poverty was examined in a retrospective study of hospital admission rate for asthma (corrected for population) and Townsend deprivation indices based on the 1991 census in the 18 districts in the West Midlands Regional Health Authority. A total of 27 117 finished consultant episodes for asthma were studied. There was a strong correlation between the hospital admission rate for asthma and the Townsend Index for each district (r = 0.82, p<0.001) (figure). The association was consistent for all age groups and remained equally strong when predominantly rural districts were excluded from the analysis. Poverty is associated with an increased incidence of hospital admissions for asthma. This is not simply due to the urban environment, suggesting that factors other than exposure to pollution are important.

Quality of life assessment in asthma patients: development and validation of a questionnaire

DT BROWN, P GRANT, J PORTLOCK, E NEVILLE School of Pharmacy and Biomedical Sciences, University of Portsmouth, Portsmouth; Chest Clinic, St Mary's Hospital, Portsmouth Health related quality of life (QOL) is a concept which is gaining increasing importance in the management of chronic diseases such as asthma. A 20 item questionnaire was designed with Likert scale questions on five different life dimensions: physical activities, role activities, emotions, social activities and health perceptions. Positive and negative items were balanced within each dimension. It was administered by interview to 70 asthma patients (35 hospital clinic and 35 community clinic) with varying severity of disease as judged by clinical criteria and 69 healthy adults from the general population, matched for age, gender and socioeconomic status. Internal consistency for the questionnaire was demonstrated by a high value of Cronbach's alpha (alpha = 0.94, n=139) and by statistically significant dimension-total correlations. Product moment correlations between the physical activities dimension score and physical measures of asthma severity (peak flow, forced vital capacity, and forced expiratory volume in one second) were statistically, highly significant. The control group established a calibration profile of typical QOL scores for the general population. Mean QOL scores for
the control group was 84-37% compared with 81-99% for mild, 67-22% for moderate and 40-80% for severe asthmatics. The questionnaire is capable of rapid administration by a physician or asthma care nurse, provides an overall QOL score which is easy to calculate and interpret for individual patients, and discriminates between patients with differing disease severity.

School absence and asthma: time for reappraisal

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Departments of General Practice, Respiratory Medicine and Epidemiology, University of Dundee, Dundee

Asthma was regarded as a major cause of school absence in the 1970s and 1980s. A follow up, for up to three school terms, of a cohort of children drawn from 12 rural, town and urban general practices in Tayside gave the opportunity to reappraise absence and asthma and the potential for a general practitioner and secondary school health worker to take anti-asthma treatment and/or smoking control (British Thoracic Society guidelines) (n=351) primary and secondary school children, taking anti-asthma treatment (British Thoracic Society guidelines) were compared with a group of non-asthmatic children, matched for age, sex and school class, to measure the effect of asthma on school absence. Working through the Education Department class registers for individual schools were examined and the number of half days absent and episodes of absence were recorded for children in the two groups. Subgroups of boys and girls, and primary and secondary school subjects were compared. The study group was also split into subgroups representing each step of asthma treatment and comparisons made between these groups. There was no significant increase in school absence, either half days missed or episodes of absence, for the children on anti-asthma treatment when compared with the matched controls. There was no difference in the amount of school absence dependent on severity of asthma as defined by the British Thoracic Society guidelines. In the early 1990s treatment of asthma, classified by the British Thoracic Society treatment steps, would not seem to be a major cause of school absence in its own right. Children with asthma who miss school are likely to have another reason for their absence.

Care of children with asthma in schools

K Mansfield, IJ Clague, EJ Pugh, PB Mattinson

Asthma Task Force, Co Durham Health Commission, Appleton House, Durham

The potential for the health and education services to form an alliance to address asthma management in children has been explored by a questionnaire directed to head teachers of all 296 primary and 43 secondary state schools in Co Durham during January 1994. Head teachers were asked about a school policy for asthma, the availability of inhaler medication, the adequacy of recording of asthma, and whether further training would be welcomed. Responses were received from 22 (76%) of primary and 28 (65%) of secondary schools. Just 40% of schools had a policy regarding care of children with asthma. Within primary schools 60% of schools allow children to be responsible for their own inhaler compared with all but one secondary school. Inhalers were locked away in 27% of primary schools. A third of schools had a member of staff trained to deal with asthma and 55% of primary and 71% of secondary schools had a member of staff who the head teacher felt could deal with an acute asthma attack. Less than half of primary and a third of secondary school teachers were aware of the range of inhalers available. In only 105 of 252 schools did the head teacher feel that the school nurse had input to the care of children with asthma. The majority of schools were interested in further training. The survey reinforces the findings in earlier studies which identify a need to develop school policies for asthma care and training of teachers. This could be achieved by closer collaboration between the health and education services.

Impact of structured nurse-led discharge planning on outcome in children hospitalised with acute asthma: a randomised controlled study

P Madge, JF Patton

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In a one year prospective audit of acute asthma care in children we identified a high incidence of poor outcomes including ongoing asthmatic symptoms and a readmission rate of 20%. This occurred despite over 93% of children receiving appropriate therapy with inhaled bronchodilators and oral corticosteroids. However, less than 10% of children had adequate discharge planning. In the present randomised, controlled study we evaluated the impact of a brief structured nurse-led discharge planning on subsequent asthma outcome. Children over two years during their first admission in the study period were randomly assigned to either a control or an intervention group. The intervention group received a specially designed information booklet on asthma, a review discussion session, a written management plan including a three day course on oral prednisolone, a two week follow up at a nurse-run asthma clinic, and access to telephone support. The control group received current standard medical care. Outcome was assessed by a postal questionnaire at 4-6 weeks after discharge, by reported reattendance for nebulised bronchodilator, or by readmission to hospital over the subsequent period (to date 8 months). There was improvement in night cough (p = 0.045), day time wheeze (p = 0.07), and breathlessness on exertion (p = 0.028) in the intervention group. Reattendances and readmissions had also decreased. These preliminary results provide clear evidence that structured nurse-led discharge planning can have a substantial impact on ongoing asthma morbidity in childhood.

Changes in lung function in asthmatic children undergoing general anaesthesia

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Department of Anaesthesia and Respiratory Unit, Royal Liverpool Children's Hospital, Alder Hey, Liverpool

Little is known of the effects of general anaesthesia on the lung function of asthmatic children. We studied 20 children with asthma and 20 children in whom asthma had been excluded, who were undergoing routine elective surgery. The children, aged between five and 15 years, were matched for age, sex and type of operation. Spirometry was performed preoperatively and twice postoperatively — "early" (the day of surgery) and "late" (the following morning). Continuous overnight pulse oximetry was also recorded. We found no difference in preoperative peak expiratory flow rate (PEFR) or forced expiratory volume in one second (FEV1) between the two groups. Both groups showed significant deteriorations in FEV1 and PEFR in the early postoperative period. Asthmatics showed significantly greater deterioration in FEV1 in the late postoperative period, whereas the control group did not. There were no significant differences in deterioration in lung function following anaesthesia between the two groups. Overnight pulse oximetry in asthmatics showed a greater range — reaching lower saturations — than in controls and time spent at saturations lower than 90% was markedly greater in asthmatics. We have demonstrated that well controlled children with asthma undergoing anaesthesia for elective surgery, demonstrate a deterioration in lung function and overnight oxygen saturation. While this did not cause any clinical problems in the group we studied, under less well controlled conditions adverse consequences may arise.

Five year follow up of patients involved in the 1989 Q fever outbreak

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Environmental Health, University of Birmingham; PHLS, Birmingham Heartlands Hospital; Chest Research Institute, Birmingham Heartlands Hospital, Birmingham

In 1989 one of the largest ever outbreaks of Q fever occurred in an area to the south of the Birmingham area. In all, 147 cases were recognised, the outbreak affecting predominantly men of working age. No deaths were recorded in the acute phase, but two developed cardiitis and two renal failure requiring dialysis. Five years after the outbreak a follow up study was performed by postal questionnaire. A total of 114 cases, for whom an address was known were circulated asking each to nominate a "buddy control" for a case control study. Replies were received from 93 (82%) patients, 37 of whom nominated a control. Fourteen had left the address, five had died, leaving 74 (61 men (84%)) for analysis. Mean age was 53 years (range 13-83), the sample being representative of the original group. Chronic symptoms were common with 69% complaining of joint pains, 66% chronic fatigue,
and 65% sleep disturbance. Cough (59%), irritability (54%), sweating (53%), memory disturbance (53%), and chest pain (51%) were complained of by more than half. Cough was complained of by 26/48 (54%) non-smokers and breathlessness by 24/48 (50%) non-smokers, neither significantly different from the symptom pattern in smokers. The case control results are being analysed. Of the five deaths two were due to cardiomyopathy possibly due to Q fever. These findings support findings from Australia that chronic Q fever itself is a real entity. Further investigation of this unique cohort is essential to investigate ongoing morbidity and possible mechanisms of this syndrome.

Case control study of clinical features associated with ampicillin resistant Haemophilus influenzae infection

SJOHNSON, RCP THOMPSON, JT MACFARLANE, H HOPFESMEURRY Respiratory Medicine, City Hospital, Nottingham; PHL, University Hospital, Nottingham Resistance to ampicillin amongst Haemophilus influenzae (HI) isolates in the UK has been steadily rising over the past decade, reaching over 8% in 1991. The consequences of this trend in clinical practice have yet to be studied. We performed a case control study of respiratory infection with HI comparing 23 patients with an ampicillin resistant isolate (ARI) and 34 controls with an ampicillin sensitive isolate (ASI). Isolates were identified from representative sputum specimens cultured from patients with at least three exacerbations of chronic pulmonary disease. The A history of cigarette smoking was less common with ARIs (14/23 compared with 29/34, odds ratio 0.12; CI 0.02 to 0.75). More patients with ARIs had received antibiotics (p=0.003), and particularly amoxicillin (p=0.0002), in the previous month. Coexisting patho-
gens were cultured from sputum with more ARIs (5 × S pneumoniai, 2 × S aureus, 1 × P aeruginosa) than with ASIs (3 × S pneumoniai × M catarrhalis, p=0.038). There was no difference in outcome between the two groups as judged by duration of hospital stay or death rate at three and 12 months. ARIs were associated with mixed infections, greater antibiotic exposure in the preceding month, particularly to amoxicillin and chronic supplicative lung disease. [Assistance acknowledged from Glaxo.]

Cefaclor MR versus co-amoxiclav in the treatment of acute exacerbations of chronic bronchitis in general practice

J HOSIE, C LIVANG, G LINDAY, D ROBERTS, G WILSON Department of Bacteriology, Southern General Hospital, Glasgow; Lilly Industries Ltd, Basingstoke Most antibiotics for the treatment of acute exacerbations of chronic bronchitis are given for seven days, although there is a trend towards five days of treatment. The common view is that patients do not receive treatment when they feel better. We describe a study where patients were allowed to take antibiotics for up to 14 days and discontinue at their own discretion. A total of 223 patients (125 men) presenting with acute exacerbation of chronic bronchitis were randomly assigned to either cefaclor MR twice daily or co-amoxiclav three times per day. Each patient was instructed to continue their treatment until they felt better. Only patients treated for more than three days were included in the analysis (112 men). Seventy cefaclor MR patients were classified as cured by their GP and 86 co-amoxiclav patients. Duration of treatment of cured patients was not significantly different, with cefaclor MR being taken for 11.4 (3.1) days and co-amoxiclav for 10.7 (3.5) days. Six cefaclor MR patients (5.9%) withdrew from the study due to adverse events compared with 13 (12.9%) of the co-amoxiclav patients (p=0.092). Patients classified as cured were interviewed one month after cessation of treatment. There was no significant difference between the rate of relapse in either treatment group. This study suggests that, contrary to popular belief, patients may require longer courses of therapy than the commonly prescribed seven or five days for their symptoms to be reduced to their satisfaction.

Impact of bronchoalveolar lavage in suspected pneumonia after renal transplantation

MJ JREDALE, SA EVANS, C BELL, MA WOODHEAD Department of Respiratory Medicine, Manchester Royal Infirmary, Manchester Bronchoalveolar lavage (BAL) is an effective means of diagnosing pulmonary infection in patients after renal transplantation. The impact of BAL on outcome in this group of patients has seldom been evaluated. We retrospectively studied all patients who had undergone BAL for suspected pneumonia after renal transplantation from January 1989 to August 1994 to evaluate whether BAL influenced therapy and/or survival. Twenty six lavages were performed on 25 patients (median age 46 years, median time from transplant 2-8 months). All patients were treated with immunosuppressive agents: 21 cyclosporin, 24 steroids, 13 azathioprine, and five antithymocyte globulin or equivalent at the time of bronchoscopy. Eighteen (69%) lavages were positive; seven for P carinii, seven for cytomegalovirus, three for fungi, two for M tuberculosis, and two for influenza A. Ten of the 18 patients with a positive BAL survived the acute episode compared with seven of eight patients with a negative BAL (p=0.11). Fifteen of the 18 patients with a positive BAL had an abnormal chest radiograph compared with one of the eight patients with a negative BAL (p=0.001). BAL findings resulted in a change in treatment in 14 cases. In 10 cases specific treatment was started. Eight (80%) of these patients survived. In four cases treatment was stopped. In four cases the BAL result led to continuation of previous empirical therapy, two (50%) of these patients survived. All patients who survived the acute episode are still alive. Median follow up for those surviving the acute episode is 20 months. In conclusion, BAL leads to a change of treatment, or confirms empirical treatment, in the majority of patients studied with beneficial effects on survival.

Comparison of pentamidine absorption in the isolated perfused rabbit lung when administered by intratracheal instillation and ultrasonic nebuliser

A BELLO, AJ HUTT, CMARROWIT Department of Pharmacy, King’s College London Pentamidine isethionate (PI) has been commonly ad-
ministered as an aerosol for the treatment of pulmonary infection by Pneumocystis carinii. However, little is known regarding the effect of this delivery method on PI absorption from the lung. Using the recirculating isolated perfused rabbit lung (RIPRL) we have investigated the absorption of PI administered by intratracheal instillation (ITI) and ultrasonic nebuliser (UN). PI was administered at 0.5, 1.0 and 2.0 mg doses to the RIPRL by ITI and UN. The mean particle size of the aerosol was determined to be 4.0 (0.1) μm. Following administration concentrations of PI in perfusate were determined by HPLC. Delivery by both methods resulted in dose related increases in maximal perfusate concentration (Cmax) and area under the perfusate concentration time curve (AUC) (table).

<table>
<thead>
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<th>Parameter</th>
<th>Method</th>
<th>Do(s) (mg)</th>
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</thead>
<tbody>
<tr>
<td>Cmax</td>
<td>ITI</td>
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</tr>
<tr>
<td></td>
<td>UN</td>
<td>3.3 (0.3)*</td>
</tr>
<tr>
<td>AUC</td>
<td>ITI</td>
<td>2.1 (0.6)</td>
</tr>
<tr>
<td></td>
<td>UN</td>
<td>5.7 (0.7)*</td>
</tr>
</tbody>
</table>

Data are mean (SD), *p<0.01 (Student’s t test).

Comparison of ITI with the UN delivery showed increases in Cmax (1.6-2.2 fold) and AUC (1.3-2.7 fold). At the 1.0 and 2.0 mg doses there was also a significant increase (p<0.05) in the Cmax values following UN administration. The increased absorption following administration by UN may be due to increased exposure of the nebulised dose to the more absorptive, highly perfused surfaces of the lung.

Effect of nebulised recombinant DNase on mediators of inflammation in cystic fibrosis

C CULLANEY, J HAYES, O’CONNOR, MF FITZGERALD Department of Medicine, University College Dublin; St Vincent’s Hospital Dublin DNA released by degenerating neutrophils is a significant contributor to airway mucus plugging. The association between hyperviscous mucus, opportunistic bacterial infection and inflammation suggests that antimucolytic agents, in addition to reducing airway obstruction, may also attenuate the inflammation-in-
Empyema of the thorax: a four year experience from Pakistan

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Ag Khan University Hospital, Karachi, Pakistan

Empyema thoracis continues to be a common and troublesome clinical problem with significant morbidity and mortality worldwide. The aetiologies involved, concomitant diseases and prognosis regarding empyema differ from series to series. There is a difference of opinion regarding its management. The aetiologies, microbiological findings and management of 81 patients of empyema were analysed. The patients were, on average, 36-4 years old; 64% had empyema due to Pseudomonas which is the most common organism. Bronchial carcinoma (9%), and hypertension (7-4%). In 62% of patients empyema developed secondary to a bronchopulmonary infection. Other aetiologies were iatrogenic (11%), infradiaphragmatic sepsis (8-6%), idiopathic (23-5%). Cultures were positive in 66-6% cases. Among the culture positive cases a single bacteria was isolated in 74%. Multiple organisms grew on the remaining 26% positive cultures.

Mycobacterium tuberculosis was isolated from 8-6% of cases. Aerobes were isolated from 92-6% of positive cultures and anaerobes from 7-4% of the remaining positive cultures. Diagnostic thoracocentesis was done on all patients and subsequently intercostal chest tube drainage was required in 57% of patients and more aggressive surgery was performed on 22% of patients; 16% of patients were treated with serial thoracocentesis and 5% received antibiotics only. Although microbiological evidence of M tuberculosis was available in seven cases, 36-7% of patients were given antituberculous therapy because of clinical suspicion of tuberculosis. The length of the hospitalisation averaged 13 days. 12% patients died, the rest recovered; 8-6% of deaths came about as a direct result of empyema. The analysis of this case series suggests that, in developing countries like Pakistan, bronchopulmonary infection is the greater cause of empyema among which tuberculosis stands out. Case fatality ratio is very high so there is need for early detection and selection of appropriate antibiotics combined with an immediate drainage procedure.

Effect of a single nebulsed dose of frusemide on cough induced by low chloride solution in patients with chronic dry cough of unknown aetiology

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Nebulsed frusemide inhibits cough induced by low chloride content solutions (LCC) in healthy subjects. We examined the effects of a single nebulsed dose of frusemide on cough induced by LCC in 10 selected patients (eight women) of mean (SD) age 53-2 (11-4) years with a chronic dry cough of unknown aetiology in a double blind placebo (saline) controlled crossover design study. These patients had a cough for >2 years and were fully investigated for a cause for the cough and had failed to respond to a course of high dose inhaled steroids and had an enhanced cough response to LCC. None were on regular medications for cough and all antiinflammatory agents were stopped 24 hours prior to which frusemide (3-75 mg/ml) in saline or saline was administered for 8 minutes via a De Vilbiss nebuliser with an output of 1 ml/min. Prior to each of the two study days, which were 1 week apart, the patients had an LCC cough challenge to determine the highest LCC which produced eight coughs in 5 minutes or more. The LCC challenge was repeated at 20, 120, and 240 minutes following frusemide or saline administration. The baseline coughs were reproducible (p=0.37). Frusemide had a weak effect on the cough response to LCC at 20 minutes (p=0.09) and 240 minutes (p=0.09) compared with saline (p=0.10 and p=0.64 respectively) (Wilcoxon signed ranks of paired individual data). Four patients showed a significant response to frusemide which was sustained up to 240 minutes (p=0.006). The weak response could be explained by hyperalgesia of the peripheral cough receptors in this select group of patients and warrants further study with chronic dosing. There is also a subgroup of "responders" who may benefit from long term frusemide.

Capsaicin cough challenge: relationship to airways obstruction and reactivity in asthma

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The capsaicin cough challenge test has previously been used to assess the sensitivity of the cough reflex in humans. However, the relationship of this test to other measures of airway function and reactivity in asthmatics is not known. The means of disease severity used were peak flow variability, % predicted FEV1, reported symptoms of cough, and the degree of non-specific bronchial hyperreactivity. We studied 43 asthmatics (27 men, mean age 49 years, range 22-73, mean (SE) % predicted FEV1, 73 (3%), mean % predicted PEF 87 (4%)]. Thirty four subjects kept a record of their peak flow and daily symptoms of cough for two weeks; 27 complained of cough more than half the days. All 43 subjects had spirometry before a capsaicin cough test which was performed using a single inhalation technique and nine increasing concentrations of capsaicin (1-500 μM). The concentration at which the subject coughed five times was noted (C5). Ten minutes after the cough test and after checking that the FEV1 had returned to baseline, we performed a histamine challenge test. The 34 subjects who kept the diary card did not differ in any variable from the remaining nine subjects. The median C5 response was the sixth concentration whilst the geometric mean lay between the fourth and fifth concentrations; this is significantly lower than normal subjects in whom the median is above the ninth concentration. The median and geometric mean PC20 for histamine were 1 (0-46) μm and the mean peak flow variability was 15 (1-36)%. Although histamine C5 responses were reduced, we found no correlation between capsaicin sensitivity (C5) and % predicted FEV1 (Pearson's r=-0.08), peak flow variability (r=0.05), or PC20 for histamine (Spearman's r=0.23). There was a weak relationship between sensitivity to capsaicin and reported symptoms of cough analysed non-parametrically (r=-0.33). In conclusion, we found no evidence that the sensitivity of the cough reflex as measured by capsaicin was related to airway calibre or bronchial hyperreactivity, suggesting that different mechanisms may be involved in the enhanced cough response of asthmatics.

Bromchodilator responsiveness of impulse oscillation model parameters in healthy and asthmatic subjects

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University Medicine, Southampton General Hospital, Southampton

Oscillometry examines the behaviour of the total respiratory system with time and frequency. Using the new Impulse Oscillation System (IOS, Jaeger, Germany) measurement function was examined in six asthmatic (A) and 10 healthy (H) subjects (mean age 37 (13) years) undergoing bromchodilator challenge (salbutamol, volumetric device). The impulse length was 1 sec to calculate central and peripheral resistance (Rz and Rp), total resistance (Rrs), lung and bronchial compliance (Cl and Cb), and central airway inerance (L2). The results are shown in the table in kPa/s.
Perception of asthma and breathlessness during exercise scored on a visual analogue scale

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Respiratory Department, Bristol Royal Infirmary, Bristol

We wished to determine whether asthmatics could distinguish between normal breathlessness and the sensation of asthma during exercise using a 100 mm visual analogue scale (VAS). 11 asthmatics (mean age 26-1 (2-9) years) with exercise induced asthma and 10 normal subjects (mean age 26-5 (5-3) years) were studied. Each performed a Balke treadmill test at 5.5 kph with a one minute incremental grade of 2% per minute to 20%, inspiring dry air at room temperature. Expired minute ventilation (Ve) was averaged over the last 30 seconds of each minute. Prior to each increase in grade the subjects quantified their sensation of breathlessness (SoB) and asthma using the VAS (Harty et al. Clin Sci 1993:85:229-36).

The relation of VAS to Ve was obtained by linear regression analysis. VAS (mm) at a Ve of 50 l/min (VAS50) and at 60% maximum Ve (VAS60) were calculated. All subjects achieved a cardiac frequency of >80% maximum predicted (range 82-104%). For all subjects there were significant linear relationships of VAS to Ve for SoB and asthma. The results are given in the table as mean (range). All indices were significantly different (p<0.05) when comparing the sensations of breathlessness and of asthma during exercise in the asthmatic subjects. We conclude that asthmatics can distinguish the sensation of asthma from that of normal breathlessness.

Comparison of two methods of collecting respiratory diary card data: conventional paper and electronic diary

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Respiratory Unit, Western General Hospital, Astra Clinical Research Unit

The study compared the suitability of conventional paper and electronic diaries (Apple Newton MessagePad with customised software) in collecting respiratory trial data. An open randomised crossover study was performed in 22 patients (13 men, age 18-70) with airflow obstruction who recorded twice daily peak flow measurements, asthma symptoms, and rescue inhaler usage in a paper (PD) and an electronic (ED) diary for one month each. Variables assessed were (1) completeness, timeliness and quality of data; (2) preferences of patients and staff for PD or ED; (3) differences in workload and costs; (4) speed of data throughput. Data were analysed using a standard parametric method for two period crossover after arcsine transformation and descriptive statistics. The results showed that (1) data reliability improved although there were more missing data with PD than ED (p=0.0001) due partly to a decision not to allow retrospective data entry on ED; (2) 59% of patients preferred the ED and 18% the PD. Age, gender and familiarity with technology had no marked association with the preference. Both ED and PD were judged easy to use; (3) days needed for data throughput decreased by 88% and staff time required by 84%; (4) an initial software fault suspended the study. Later hardware power faults caused problems. New versions appear to have overcome these. We conclude that (1) ED offers potential benefits of improved data quality, faster data throughput, and results in possible cost savings; (2) there is a high degree of patient acceptability for the ED.

Respiratory observations on trekkers accompanying the 40th anniversary British Everest Expedition

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Department of Anaesthetics, University Hospital of Wales, Cardiff; Department of Respiratory Medicine, Western Infirmary, Glasgow; Newcastle Medical School

Little is known of the respiratory parameters of subjects as they acclimatise to altitude. We have measured FVC, SpO2 and PAO2 on 39 trekkers as they walked from Jiri (1860 m) to Base Camp Everest (5550 m) over a three week period. PambO2 was predicted from known altitudes (Kuntz et al. 1906; West et al. 1983) and used to calibrate a fuel cell which measured PAO2 following a maximum expiratory manoeuvre. Percutaneous oxygen saturation (SpO2) was measured using a pulse oximeter (PneuPAC). Oxygen saturation (SaO2(calc)) was also predicted assuming PaO2 equal to the PAO2 using an algorithm (Mohan et al. 1977) in which predicted PaCO2 (RQ=0.85) and arterial pH are factors. It has been shown that pH changes little below 5000 m (Dill et al. 1937; West et al. 1962). The results demonstrate the progress of acclimatisation in the face of increasing altitude. The observed SpO2 was increasingly unstable above 3440 m. This suggests instability either of ventilation or of V/Q matching. The discrepancy between measured SpO2 and predicted SaO2 (calc) (p<0.0001 for the higher altitudes) may indicate a deterioration in pulmonary function caused by pulmonary oedema.

Comparison of respiratory function after single lung transplantation for fibrosing alveolitis and emphysema

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Departments of Respiratory Medicine and Cardiothoracic Surgery, Freeman Hospital, Newcastle upon Tyne

Single lung transplantation (SLT) is performed in patients with both advanced fibrosing alveolitis (CFA) and emphysema (E). We have compared 10 patients receiving SLT for E with six receiving SLT for CFA to assess how the native lung influences tests of overall respiratory function. Patients were studied for a mean of 17 months (range 6-48) after SLT when free of complications. None had had evidence of obliterative bronchiolitis. Investigations included lung volumes (by plethysmography), spirometry, MEVF curves (Vemax and moments analysis), TLC, and Kco and quantitative V/Q scans using 133Xe and 99mTc respectively. Relative V was
measured after a single full inspiration. The results are shown in the
tables. In comparison with patients receiving SLT for CFA, those
tidal flow-time or flow-volume tracings (Morris et al. ERJ 1990;3:
901-9). The aim of this study was to assess further the validity of
this measurement. During tidal breathing EV was measured in two
groups of subjects: (1) seven normal subjects in whom, during relaxed
tidal breathing, two expiratory resistances of increasing severity were
assessed in two different settings; (2) eight patients with emphysema
were measured by pneumotachography. Paired r tests showed that
there was no difference between the EV increments measured by
the two methods (low resistance – spirometric volume increment
(SVI) = 0.271 (mean (SD)), EV = 0.23 (0.08); higher resistance – SVI
= 0.65 (0.18), EV = 0.52 (0.09)). Regression analysis showed that
the lower resistance, SVI was significantly related to EV, SVI =
0.003 + 1.15 EV, r² = 0.77, p<0.01. With the higher resistance the
regression relationship between these two variables was not significant.
(2) Seventy nine patients with a clinical diagnosis of airflow obstruction
(AFO) in whom thoracic gas volume was measured by constant
volume plethysmography. In this group of patients (FRC % predicted
= 153 (44)), EV predicted the degree of overinflation according to
the equation, FRC % predicted = 110 + 149 EV, r² = 0.46, p<0.001. It is
hypothesised that EV measures the volume that EEV is above the
relaxation volume of the respiratory system for that part of the lung
with open airways over the tidal volume range, and thus does not
include the volume increment caused by trapped gas.

with emphysema have similar perfusion but relatively more ventilation
of the transplanted lung and better overall CO transfer, despite
generally worse airway function. The larger TLC and MEFV moments
are attributable to the volume and very slow emptying of the native
emphysematous lung. Tests of airway function are likely to be less
sensitive for detecting complications after SLT for emphysema
compared with CFA.

<p>| Table 1 Mean (SE) lung volumes, CO transfer and scans |
|-------------|-------------|-------------|-------------|-------------|-------------|</p>
<table>
<thead>
<tr>
<th>TLC (L)</th>
<th>VC (L)</th>
<th>TLC (%)</th>
<th>KCO (mmol/L/min)</th>
<th>Graft Q (%)</th>
<th>Graft V (%)</th>
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<tr>
<td>CFA</td>
<td>103</td>
<td>76-3</td>
<td>56-7</td>
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</tr>
<tr>
<td>E</td>
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<tr>
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| Table 2 Mean (SE) MEFV curves |
|-----------------------------|-----------------------------|-----------------------------|
| Vmax (L) | Moments |
|-------------|-------------|-------------|-------------|-------------|-------------|
| 75%VVC | 50%VVC | 25%VVC |
|-------------|-------------|-------------|-------------|-------------|-------------|
| CFA       | 84-0        | 57-0        | 46-8        | 0.72        | 1.07        | 2.66        |
| E         | 56-5        | 40-5        | 34-2        | 0.94        | 2.25        | 7.95        |
| p         | 0.046 NS    | 0.035 NS    | 0.013        | 0.014       |             |             |

Time constants during tidal expiration and maximally forced expiration
F DENBY, KG MADGWICK, MJ MORRIS Osher Chest Unit, Churchill Hos-
pital, Oxford From a maximally forced flow volume curve from total
lung capacity (TLC) the time constant, ts, of the respiratory system
can be calculated as the ratio of volume/flow. It has been shown that
the time constant of a fully relaxed expiration from TLC is not different
from that of a maximally forced expiration. We have postulated that
the time constant of the respiratory system can be estimated from
analysis of the last part of tidal expiration when relaxation of inspiratory
muscles has occurred. The aim of this study was to compare time
constants measured from tidal expiration, tsTID, in normal subjects
and in patients with airflow obstruction (AFO) with the values
obtained from maximally forced flow volume curves with volume
measured (a) by integrating flow at the mouth, trsMO, and (b) at the
chest wall using transmural body plethysmography, trsCH. In 10
normal subjects (normal (SD) PEV, % predicted was 120 (16) and in
11 patients with AFO PEV, % predicted was 48 (21). There was a
significant difference between the normal subjects and the AFO
patients in the time constants measured by each of the three methods.

<table>
<thead>
<tr>
<th>tsTID(s)</th>
<th>trsMO(s)</th>
<th>trsCH(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal subjects</td>
<td>0.45 (0-07)</td>
<td>0.48 (0-13)</td>
</tr>
<tr>
<td>AFO patients</td>
<td>1.5 (1-04)</td>
<td>3-1 (3-3)</td>
</tr>
</tbody>
</table>

In all the subjects the difference between trsMO and trsCH was
significantly related to dv, the maximum volume of alveolar gas
compression, difference = 1.55 + 4.3 dv, r² = 0.57, p<0.001. In the
normal subjects there was no important difference between ts meas-
ured in all three ways; however, in the AFO patients tsTID was closer
to trsCH than to trsMO, probably because alveolar gas compression is
not occurring during tidal expiration.

Evaluation of an index of overinflation derived from analysis of
tidal breathing
E GRINT, KG MADGWICK, MJ MORRIS Osher Chest Unit, Churchill Hos-
pital, Oxford An index of overinflation (EV) can be derived from

Increased QT dispersion: a novel marker of abnormal myo-
cardial repolarisation during acute hypoxaemia
DG KIELY, RG CARGILL, AD STRUTHERS, BJ LIPWORTH Department of
Clinical Pharmacology, Ninewells Hospital and Medical School,
Dundee Prolongation of QT interval has been associated with cardiac
dysrhythmias and sudden death, reflecting abnormal myocardial re-
porlarisation. Hypoxaemia and β agonists have both previously
been shown to increase QT interval, raising the possibility of deleterious
synergistic effects in acute severe asthma. Recently QT dispersion
(interlead variability in QT interval) has been proposed as being more
sensitive than QT interval as a marker of repolarisation abnormalities.
We have therefore evaluated the effects of acute hypoxaemia on both
QT interval and QT dispersion in eight normal men. After resting to
achieve baseline haemodynamics, subjects were rendered hypoxaemic
for 30 minutes by breathing an N₂O mixture to achieve Sao₂ of
75-80%. From the ECG, lead II corrected QT interval (QTc) and over-
all corrected QT dispersion were measured using a computer
linked digitising tablet according to standard criteria. QTc dispersion
was significantly increased during hypoxaemia compared with baseline
at 69 (6) v 50 (5) ms respectively (p<0.05), 95% CI for mean
difference 2-35 ms, whilst QTc interval was not significantly affected
by hypoxaemia: 428 (8) v 416 (10) ms (p = 0.3). This was not
associated with any change in serum potassium: 4-07 (0-06) v 4-06
(0-07) mmol/l (p = 0.62). Plasma catecholamines were not sig-
nificantly affected by hypoxaemia: noradrenaline 4-75 (0-65) v 3-97
(0-36) mmol/l (p = 0.11), adrenaline 0-22 (0-027) v 0-16 (0-06) mmol/
l (p = 0.15). There were significant increases in both heart rate (HR) and
mean pulmonary arterial pressure (MPAP) in response to hyp-
oxiaemia: HR 78 (3-4) v 64 (1-9) bpm (p<0.05) and MPAP 23 (1-2)
v 9 (0-9) mmHg (p<0.05). In summary, QTc dispersion was found to be
a more sensitive marker of repolarisation abnormalities than
QTc interval during acute hypoxaemia. Furthermore, significant in-
creases in QTc dispersion cannot be explained by changes in cir-
culating catecholamines. Thus, measurement of QTc dispersion may
be a useful clinical tool for predicting patients at risk of hypoxaemia
associated arrhythmias.

Placebo controlled study comparing budesonide 256 μg and
128 μg with fluticasone propionate 200 μg in patients with sea-
sonal allergic rhinitis
MA STEARN, R DAHL, C SCHEREVIELS Midlands Asthma and Allergy
Research Association (MAARA), Leicestershire General Hospital, Leicester,
UK; Aarhus Kommunehospital, 8000 Aarhus C, Denmark; Astra Draco
AB, Clinical Research and Development, Lund, Sweden In a two centre
study in the UK and Denmark 602 patients (≥18 years) were eval-
uated (according to all patient treated analysis) to compare the
efficacy and safety of budesonide aqueous nasal spray 256 μg
(BUD256), 128 μg (BUD128), placebo and fluticasone propionate
aqueous nasal spray 200 μg (FP200), all once daily, in the treatment
of seasonal allergic rhinitis. We used a parallel, randomised design,
double blind for BUD and placebo and single blind to the investigator
for FP. A one week run in period preceded trial medication for 4–6 weeks during the grass pollen season. Patients recorded nasal symptoms on a Diary Card. All active treatments produced significantly lower mean scores than did placebo for all nasal symptoms (p<0.05). BUD256 was significantly better than FP200 in reducing sneezing (p<0.05). We found no differences in effects between BUD128 and FP200. Adverse events recorded during study were mild and transient for each treatment group. The results suggest that the clinical effect of BUD128 and FP in the nose is similar and that BUD256 is more effective in seasonal allergic rhinitis than FP200.

Study to compare the efficacy of budesonide (Pulmicort® Turbuhaler®) and fluticasone propionate (Flixotide® Diskhaler®) in the treatment of asthma

G BASRAN, R SCOTT, M CAMPBELL, A KNOX, R SMITH, J VERNON, A WADE for a UK STUDY WORKING GROUP Rotherham District Hospital, Moorgate Road, Rotherham The study objective was to compare the efficacy of budesonide Turbuhaler (Bud TBH) with that of fluticasone propionate Diskhaler (FP DH), at half the baseline dose, in asthmatic patients requiring daily doses of 400 or 800 µg of conventional inhaled steroids. Only patients with symptomatic asthma during run in were randomised. The study was of a multicentre, open, parallel group design with treatment for eight weeks with Bud TBH or FP DH at doses of 100 or 200 µg bid. Data were analysed by an all patients treated (APT) approach. Change from baseline in morning PEF was the primary variable. Secondary variables included asthma symptom score, β₂ agonist use, FEV₁, and FVC. Treatments were compared by ANOVA. One hundred and seventy six patients were randomised and 171 (78.93 M:F; mean age 39.9 years) were valid for APT analysis. The mean morning PEF during run in was 408.9 (Bud TBH) and 408.5 min (FP DH). There were no clinically or statistically significant differences between the treatments in any of the variables recorded in diary cards or at clinic visits. Mean increase in morning PEF (adjusted for age and sex) was 8.0 (Bud TBH) and 16.6 (FP DH) l/min (NS, p=0.33). This study does not show any differences in efficacy between Bud Turbuhaler and FP Diskhaler when halving the baseline dose of conventional steroids.

Ipratropium bromide given by Turbuhaler® is more potent than when given by pressurised metered dose inhaler (MDI)

SP MATUSIEWICZ, FGE BÖLLERT, M DEWAR, G BROWN, A MCLEAN, AP GREENING, GK CROMPTON Respiratory Unit, Western General Hospital, Astra Clinical Research Unit, Edinburgh A study was undertaken to determine the relative potency of ipratropium bromide (IBPb) via Turbuhaler in comparison with pressurised metered dose inhaler (MDI) in patients with reversible airflow obstruction. The design of the study was randomised, double blind, double dummy placebo controlled, single dose crossover study with one screening visit and five study visits. FEV₁ was recorded five minutes before and 5, 15, 30, 60, 90, 120, 180, 240, 300, 360 minutes after drug administration. The end points of FEV₁ and area under the curve (AUC) analysis were examined by analysis of variance. Thirty three patients (21 men, mean age 55.6 years with reversible airflow obstruction (FEV₁ 35–80% of predicted and ≥1 litre demonstrating ≥15% reversibility to IBPb) completed the study. The % change in FEV₁, is shown in the figure. In clinically stable reversible airflow obstruction ipratropium bromide via Turbuhaler elicited a dose dependent increase in FEV₁, which peaked at 20–40 µg. AUC analysis showed that 20 µg via Turbuhaler was more effective than 20 µg via MDI. The 20 µg MDI was not different from 10 µg Turbuhaler. The efficacy ratio of Turbuhaler: MDI was round 1:5–0/2 and is similar to that demonstrated with other drugs administered by Turbuhaler.

Comparative efficacy of ipratropium bromide via Turbuhaler® and MDI in patients with reversible airflow obstruction

FG E BÖLLERT, SP MATUSIEWICZ, M DEWAR, G BROWN, A MCLEAN, AP GREENING, GK CROMPTON Respiratory Unit, Western General Hospital, Astra Clinical Research Unit, Edinburgh Previous studies with other drugs have shown Turbuhaler (TH) to be more effective than pressurised metered dose inhaler (MDI) (Thorax 1993:48:434; Chest 1994;105:697–700). We therefore aimed to compare the efficacy of ipratropium bromide (IB) via TH and MDI. A randomised, double blind, double dummy, crossover dose study was performed. A total dose of 160 µg IB was given as individual doses of 20, 20, 40 and 80 µg via TH or MDI at 45 minute intervals on two study days at least 48 hours apart. FEV₁ was measured prior to the first dose and 40 minutes after dosing. Analysis of variance for the end point PEFR was performed. Fifteen patients (nine women, mean age 60 years, mean baseline FEV₁, 1.521, 57% of predicted normal with >15% reversibility to 40 µg IB) completed the study. There was a steep dose-dependent increase in FEV₁, irrespective of the inhaler device, 9/15 patients on TH and 5/15 on MDI reached the maximum response at or before the cumulative dose of 40 µg IB.

Mean (SD) cumulative dose response to IB via TH and MDI and mean ratio (% of TH:MDI FEV₁ response)

<table>
<thead>
<tr>
<th>Cum. doses</th>
<th>FEV₁ response</th>
<th>Mean ratio (% adjusted for period and baseline FEV₁ (SD))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Base</td>
<td>1.5 (0.5)</td>
<td>1.46 (0.47) N/A</td>
</tr>
<tr>
<td>20 µg IB</td>
<td>1.79 (0.6)</td>
<td>1.71 (0.54) 102.9 (98.8–107.1), p=0.16</td>
</tr>
<tr>
<td>40 µg IB</td>
<td>1.88 (0.65)</td>
<td>1.82 (0.59) 100.8 (96-1–105.8), p=0.71</td>
</tr>
</tbody>
</table>

There was no significant difference between the devices at any dose of IB. We conclude that there was no significant difference in the efficacy of TH and MDI at the dose range of IB tested, but further studies to assess the relationship at lower doses of IB might be warranted. A proportion of asthmatic patients respond to low doses of IB.

Comparison of the efficacy and tolerability of efmorotol and salbutamol dry powder capsules for inhalation in elderly asthmatic patients

RM ANGUS, NC THOMSON for a UK EFORMOTEROL STUDY GROUP Department of Respiratory Medicine, Western Infirmary, Glasgow Eformotol is a selective β₂ agonist which when given by inhalation produces bronchodilation of rapid onset which lasts for 12 hours. The aim of this study was to examine the efficacy and tolerability of eformotol dry powder in comparison with salbutamol dry powder in elderly asthmatic patients. A multicentre, parallel group design was employed. Two hundred and sixty two patients (105 women) of mean (SD) age 71 (4) years, range 64–82 years, were studied. All had demonstrated at least 15% reversibility to inhaled salbutamol. Patients were randomised to receive either eformotol 12.5 µg bd, eformotol 24 µg bd, or salbutamol 400 µg qid for a three month period. During a two week run in patients received salbutamol 400 µg qid then entered one of the treatment arms. Sixty four patients were withdrawn during the study: 21 (25%)
efomterol 12 µg bd, 16 (18%) efomterol 24 µg bd, and 27 (31%) salbutamol 400 µg qid. Efomterol 24 µg and 12 µg were statistically significantly different from salbutamol: the overall mean difference in the mean efficacy variable, pretreatment morning PEFR recorded daily by the patient, was estimated to be 33 and 34 m/min for efomterol 12 µg and 24 µg, respectively. Moreover, the mean pre-treatment evening PEFR for efomterol were both different from salbutamol by 32 and 30 m/min respectively. The daily use of rescue medication was less in the 24 µg efomterol group when compared with salbutamol. Analysis of asthma scores and sleep disturbance did not show a significant difference, however the patients' overall assessment of the treatment was significantly better in both the efomterol groups when compared with salbutamol. Laboratory abnormalities did not seem to follow any particular pattern within the treatment groups. Similarly, there were no clinically significant changes in pulse, BP or ECG measurements. Overall tolerability was assessed by patients as good or very good for efomterol 12 µg (89%) and 24 µg (88%) compared with (82%) for salbutamol (NS). One patient in each of the 12 µg efomterol and salbutamol group had a serious adverse event that may have been drug related. These results suggest that efomterol 12 µg and 24 µg bd provide superior asthma control and are tolerated at least as well as salbutamol 400 µg qid in the elderly.

Regular formoterol treatment in mild asthma: effect on bronchial reactivity during and after treatment

DH YATES, MW WORSDELL, PJ BARNES, KP CHUNG Department of Thoracic Medicine, National Heart and Lung Institute and Royal Brompton Hospital, London There has been concern regarding the potential detrimental effects of β2 adrenoceptor agonists on bronchial responsiveness and development of tachyphylaxis to their protective effects against bronchoconstriction in asthma. Formoterol, a long-acting β2 agonist, is effective in single doses in the prevention of methacholine-induced bronchoconstriction. In a double blind, placebo controlled crossover study we examined the effect of regular formoterol treatment (24 µg twice daily over two weeks) on airway calibre and bronchial responsiveness to FEV1 and bronchial reactivity to methacholine were also measured after treatment cessation at 12, 36, 60 and 108 hours and at two weeks in order to examine for a “rebound” increase in BHR. FEV1 and PC20 rose significantly with formoterol (3-4 to 3-79 l; and 0-53 mg/ml to 2 mg/ml; p<0-005 compared with placebo) measured 12 hours after one dose. This effect was not maintained after two weeks treatment, FEV1, falling to 3-51 l and PC20 to 0-93 mg/ml (p<0-007 and p<0-04 respectively compared with placebo), but both remained significantly higher than baseline. No significant rebound decreased in FEV1 was seen after the formoterol withdrawal. Significant tachyphylaxis to both the bronchodilator and broncho-protective effects of formoterol thus occurred. There was no evidence of rebound BHR. Regular usage of formoterol may therefore be accompanied by partial loss of its beneficial effects. The clinical significance of these findings remains to be determined. [We thank C. P. C. Geigy, Switzerland for supply of formoterol and for financial support.]

Efficacy of once daily budesonide via Turbohaler® in moderate asthma

LM CAMPBELL ON BEHALF OF THE PATRON INVESTIGATORS GROUP Southbank Surgery, Kirkintilloch, Glasgow Once daily budesonide has been shown to be as effective as twice daily in mild asthma (Jones et al. Respir Med 1994;88:293–6). We undertook this study to test the concept of once daily inhaled budesonide in the control of more severe asthma. Patients who were receiving 400–600 µg inhaled steroid daily entered a one week run in to assess peak expiratory flow rate (PEFR), asthma symptoms, and rescue β2 agonist use. One hundred and sixty four symptomatic patients were randomised to open, parallel group treatment with 800 µg budesonide via Pulmicort® Turbohaler® administered either as 800 µg once daily (n=109) or as 400 µg twice daily. PEFR and symptoms were reviewed at clinic visits at four and eight weeks, and patients kept a daily record of symptoms, PEFR (am and pm) and β2 agonist use. Morning PEFR recorded in diary cards increased by 21 l/min in the twice daily group (p<0-0001) and by 29 l/min in the once daily group by eight weeks (p<0-001), with overall improvement in symptom scores in both groups. Diary card records also showed significant improvements in PEFR and reduction in diurnal variation, with reduced β2 agonist use. Day time and night time symptoms were reduced in both groups. There were no significant differences found between the treatment groups for any of the variables tested. We conclude that 800 µg budesonide can be given equally effectively either as a once daily or twice daily dose via Turbohaler.

RFLP analysis of the P2 gene of Haemophilus influenzae isolates recovered from patients with bronchietasis

A PYE, RA STOCKLEY, TF MURPHY, SL HILL Lung Immunobiological Research Laboratory, The General Hospital, Birmingham; Suny at Buffalo, Buffalo, USA P2 is the major outer membrane protein of non-typable Haemophilus influenzae isolated from bronchietatic patients. This protein has been shown to express a highly strain specific immunodominant epitope. Phenotypic changes have been demonstrated in this protein in isolates recovered longitudinally from patients with bronchietasis. Such changes in P2 could provide a mechanism for NTHI to evade lung host defence systems thereby allowing its persistence in the lower respiratory tract. We report the preliminary results of RFLP analysis of the P2 gene in 52 isolates of NTHI collected from 10 patients with bronchietasis. RFLP analysis was performed on a PCR amplified gene segment (using appropriate oligonucleotide strain) and the restriction endonuclease XbaI (which has three recognition sites in this segment of the reference strain). None of the 52 isolates (eight from one patient) had no XbaI sites anywhere in the amplified segment. The restriction patterns also revealed that none of the isolates had the XbaI restriction site in the variable region of loop 4. In the remaining isolates a variety of restriction patterns were observed for either or both of the two other restriction sites in conserved regions. In the 10 individual patients, RFLP analysis showed that three of four patients' isolates collected sequentially showed different P2 restriction patterns even though P2 was phenotypically similar on SDS-PAGE. In five of the remaining six patients where the phenotype of P2 differed on SDS-PAGE in isolates collected sequentially there were corresponding differences in RFLP. In conclusion, changes in the P2 gene occur more frequently in NTHI isolates from patients with bronchietasis than have previously been suggested, even in the presence of apparently similar phenotype on SDS. Further studies are required to relate these changes to the expression of relevant antigenic epitopes.

An air interface organ culture model for use in the study of the interactions of bacteria, viruses and pharmacological agents with the human respiratory mucosa

AD JACKSON, GF RAYNER, A DEWAR, PJ COLE, R WILSON Host Defence Unit, Department of Thoracic Medicine, Royal Brompton National Heart and Lung Institute, London The mucociliary epithelium lining the respiratory tract is the primary interface between airborne particles, vapours and gases, and the host epithelium. Respiratory organ cultures that are immersed in culture medium are unphysiological. We have assessed a human respiratory tissue organ culture model with an air interface by light microscopy, scanning electron microscopy (SEM), and transmission electron microscopy (TEM). Dissected nasal turbinate tissue was soaked in antibiotic medium to remove commensal flora, and was maintained in a humidified atmosphere with 5% CO2 on a filter paper strip kept whose ends were immersed in medium. Without changing the culture medium ciliary beat frequency (CBF) remained constant (11–12 Hz) over five days but fell significantly (p=0.004) at 10 days to 7-9 (0.8) Hz. By SEM, percentage tissue cover by mucus varied from 29–64% with no significant time related trend. The percent of the surface covered by ciliated cells was significantly reduced at five and 10 days. By TEM nuclear heterochromatin was reduced at five days and 10 days. Mitochondria had normal ultrastructure at four days but appeared slightly abnormal at five days and severely damaged at 10 days. Ciliary density on individual ciliated cells was significantly reduced at 10 days. Tight junctions between cells, cell projection from the organ culture surface, cytoplasmic blebbing and cell vacuolation remained normal over 10 days. When the culture medium was replaced daily TEM indicated that mitochondria and nuclei were normal, CBF was between 10 and 12 Hz and percentage tissue cover by mucus in individual cells was normal after 20 days. This model is simple to construct and reproduces physiological conditions in vitro. Because of its long viability it can be used to study virus and bacterial infections, and the effects of pharmacological agents and environmental factors.
Recognition of outer membrane proteins of non-typeable Haemophilus influenzae by IgG subclass antibodies in serum and sputum from patients with bronchiectasis

JL MITCHELL, SI HILL
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Non-typeable Haemophilus influenzae (NTHI) is frequently isolated from the sputum of patients with bronchiectasis. The organism persists in the airways of these patients despite the presence of an exuberant inflammatory response which includes a significant local production of immunoglobulin in the lung secretions. We have therefore compared the NTHI-specific IgG subclass responses in pooled sputum and serum collected from six patients with bronchiectasis to assess the relative importance of outer membrane proteins (OMP) or NTHI in eliciting antibody responses in the lung and blood. OMP complexes from eleven subtypes (based on the relative molecular weight of the proteins P2 and P5 on SDS-PAGE) of NTHI were subjected to SDS-PAGE, blotted onto nitrocellulose and then incubated in the presence of either the serum pool, or the sputum pool pool. This was followed by incubation with biotinylated polyclonal antibodies to one of the four human IgG subclasses and colour development with HRP-conjugated avidin-biotin complexes. In addition, the outer membrane protein P6 was purified from NTHI and used alone in similar blotting experiments. The results showed recognition of all of the major OMPs of the subtypes of NTHI tested. Some antigens were more strongly recognised by one subclass of IgG than by others. Such differences were apparent in the serum and sputum pools. Of major importance was the strong recognition (particularly in sputum) of a 16 kDa antigen consistent with the OMP P6 detected by immunoblot with sputum at titres as low as 1/1250. There was no recognition of the band in the 16 kDa position after the whole cell culture was subjected to neuraminidase K digestion confirming that this result was not due to contaminating lipopolysaccharide. In addition similar results were obtained with the purified P6. These experiments suggest that P6 is an important immunogen in the airways of patients with bronchiectasis. Further studies are necessary to establish the functional role of P6 specific antibodies in the lung secretions of these patients.

Effect of erythromycin on Haemophilus influenzae endotoxin (HIE)-induced release of inflammatory mediators by human bronchial epithelial cells (HBEC) in vitro

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Department of Respiratory Medicine and Allergy, St Bartholomew’s Hospital, London
Although several studies have demonstrated that low dose, long term erythromycin (ERM) treatment is effective in the management of chronic lower respiratory tract infections such as chronic bronchitis and bronchiectasis, the mechanisms underlying the action of ERM are not clear. We have cultured HBEC as explant cultures from surgical tissue and investigated the effect of ERM on HIE-induced release of inflammatory mediators in these cultures. Confluent cultures were incubated for 24 hours with 100 μg/ml HIE and analysed for interleukin 6 and 8 (IL-6 and IL-8) and soluble intercellular adhesion molecule 1 (sICAM-1) released into the culture medium. Additionally, cultures were incubated in the presence of 0.1-10 μg/ml ERM and investigated for the effect of this antibiotic on HIE-induced release of IL-6 and IL-8. IL-6, IL-8 and sICAM-1 in the medium were analysed by ELISA. HIE significantly increased the release of IL-6 and IL-8, from 0.9 (1.5) pg IL-6/μg cellular protein and 83.7 (28) pg IL-8/μg cellular protein in control untreated cultures, to 12.12 (1.5) pg IL-6/μg cellular protein (p<0.005) and 225.7 (44) pg IL-8/μg cellular protein (p<0.005). HIE significantly increased the release of sICAM-1 from 0.23 (0.03) ng sICAM-1/μg cellular protein in control cultures, to 3.82 (0.9) ng sICAM-1/μg cellular protein (p<0.001). Incubation of HBEC in the presence of 0.1-10 μg/ml erythromycin significantly blocked the HIE-induced release of both IL-6 and IL-8. These results suggest that H influenzae-induced release of inflammatory mediators from airway epithelial cells may contribute to chronic airway inflammation and that this effect may be modulated by erythromycin treatment.

Late as well as early neutrophil influx is CD18 dependent in E coli pneumonia

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A prerequisite for resolving inflammation is cessation of neutrophil influx. A mechanism by which this might be controlled is switching of adhesion molecule dependence. Winn and Harlan (J Clin Invest 1993;92:1168-73) showed that late neutrophil influx into the peritoneum becomes CD18 independent. The requirement for CD18 differs in pulmonary and systemic circulations. Hence we studied early and late CD18 requirements and anti-CD18 effects in C57BL/6 mice infected with E coli. Pairs of rabbits had E coli instilled bronchoscopically into the right cranial lobe. After six or 30 hours they received either anti-CD18 (Celtech) or saline followed by 111In labelled neutrophils from a donor. Six and 12 hours later they received repeat antibody or saline and after 20 hours received 125I labelled red blood cells. A dual isotope gamma camera image was obtained. One hour after this the lungs were lavaged. Lavagate and tissue gamma activity were counted. The neutrophils accumulating from six or 30 hours were calculated as the iodine accummulating in excess of that in the blood pool as a percentage of injectate. The results show that neutrophil influx was significantly (though not completely) inhibited by anti-CD18 antibody, not only at early (six hour) but also at late (30 hour) time points. Lavage neutrophil accumulation was not significantly affected, perhaps because of a greater effect on sequestration than migration, or because anti-CD18 coated cells are easier to lavage out of the lungs. [Work supported by MRC (UK), Anti-CD18 supplied by Celtech.]

Analysis of lymphocyte proliferation and interleukin production in response to CMV antigen following lung transplantation

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Cardiopulmonary Transplant Unit, Department of Microbiology, Freeman Hospital, Newcastle upon Tyne; Department of Virology, Newcastale University
As part of a prospective study conducted over the last two years blood and BAL lymphocyte cultures were tested for their degree of CMV specific lymphocyte proliferation, by using a triated thymidine uptake assay. This was done at the time of routine (one week, one month, three month, six month, and yearly) and emergency diagnostic bronchoscopies. Culture supernatants were tested for IL-2 and IL-4 levels using Medgenix ELISA kits. There was no association between LB (lymphocyte proliferation blood) or LPB lymphocyte proliferation (lymphocyte proliferation lavage) positivity and CMV excretion/non-excretion or CMV antigenemia at the time of the bronchoscopy. There was a significantly higher rate of LB positivity (p=0.0015) amongst the group with a history of CMV excretion at some time than those who never excreted CMV. This suggests that the LPB positivity may predate or postdate other episodes of excretion that our surveillance and diagnostic bronchosopies did not detect. There is no association at the time of bronchoscopy between LB or LP and clinical status, rejection incidence, infection incidence, or OB. Previous work presented at the BTS has shown that asymptomatic CMV excretion is not associated with a greater incidence of infection, rejection, or problems with lung function. The finding of IL-2 or IL-4 in the supernatant following LB positivity is almost significantly greater amongst the group with a history of CMV excretion (analysis not limited to episodes of CMV excretion) compared with the group that never excreted CMV. The finding of IL-2 or IL-4 in the super- nutant following LPB or LPL positivity does not correlate, at the time of bronchoscopy, with infection, rejection, or clinical status. The three patients in the study who had CMV disease (two died, one survivor) never had a LPB or LPL response. Perhaps the exuberant/ disordered immune response in CMV disease reflects an inability of the patients’ lymphocytes to respond to CMV antigen. Patients who had histological evidence of CMV cells (cytopathological effects) but no other inflammatory changes exhibited similar LB and LPL responses to the CMV excretor group. They did not mimic the CMV disease group in consistently negative LPB and LPL. [This study was supported by British Lung Foundation.]
Effect of salmeterol on the interaction between Pseudomonas aeruginosa (PA) and the respiratory mucosa in vitro

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Host Defence Unit, Department of Thoracic Medicine, Royal Brompton National Heart and Lung Institute, London

Salmeterol is a potent acting β2 agonist which we have previously shown to reduce the ciliary beat slowing and epithelial disruption caused by the Pseudomonas aeruginosa (PA) toxin pyocyanin (Br J Pharmacol 1994;112:493-8). We have studied the effect of salmeterol on PA interaction with the respiratory mucosa of an organ culture with an air-mucosal interface by scanning electron microscopy. Human adenoid or turbinate tissue was incubated with 4 x 10^(-7) M salmeterol for 30 minutes prior to inoculation with PA (A), or 20 µl 4 x 10^(-7) M salmeterol was pipetted onto the organ culture surface immediately prior to bacterial inoculation (B). Infected organ cultures at eight hours showed a significant (p<0.02) increase in epithelial damage compared with control, and PA was predominantly associated with mucus and damaged cells. Salmeterol significantly (A: p<0.04, B: p<0.05) reduced epithelial damage caused by PA infection. Salmeterol had no effect on either the number or distribution of PA adhering to the mucosa. These results show that salmeterol reduces the damage caused by PA infection of the respiratory mucosa in vitro and may benefit patients infected by PA by this mechanism as well as bronchodilatation.

Inhibition of substance P degradation and its effect on human respiratory ciliary activity in vitro

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The tachykinin substance P (SP) is an important mediator in many inflammatory conditions of the lung. We have previously demonstrated that SP (10^(-7) to 10^(-4) M) alone and in combination with calcium ionophore A23187 (10(-9) M) induces ciliary disorientation in tracheal epithelial cells, which is associated with a transient increase in ciliary beat frequency (CBF) (Eur Respir J 1991;4(Suppl 14):291S). We hypothesised that the breakdown of SP (mediated by the membrane-bound enzyme neutral endopeptidase) was responsible for the rapid termination of this CBF response. Here we studied the CBF response to SP (10^(-4) M) in the presence of the endopeptidase inhibitor phosphoramidon (10^(-4) M). Respiratory epithelial cells, obtained by brushing the inferior turbinate of 10 normal volunteers, were suspended in medium 199 and placed in a perfusion chamber. CBF was measured using a videomicroscopy system (Respir Med 1994;88:89-101) and the rise in CBF to one of the following was observed: (1) phosphoramidon alone, (2) substance P alone (3) phosphoramidon + substance P. We found that phosphoramidon alone had no effect on baseline CBF and confirmed that SP alone produced a transient rise. However, when SP and phosphoramidon were perfused together the transient response was no longer present but was replaced by a delayed rise which was significantly elevated above control at 25 minutes (table). The time course of this delayed rise in CBF is similar to that we have previously reported with calcium ionophore A23187 (J Physiol 1991;439:103-13). Furthermore, the delayed response was abolished by the calcium channel blocker lanthanum chloride (250 µM). We conclude that inhibition of neutral endopeptidase activity modifies the time course of the SP-induced increase in CBF to produce a calcium dependent, delayed rise in CBF.

Primary and secondary ciliary disorder

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Cilia on the different cell types in the respiratory system have characteristic ultrastructure, which can be assessed by electron microscopy. Patients with ciliary disorder (CD) have impaired mucociliary clearance (MCC) because cilia on the same cell beat in different directions. CD can be measured by the standard deviation (SD) of the angles that the ciliary axles on a cell make with the perpendicular – that is, the higher the SD, the greater the CD. CD may occur secondary to viral infection or chronic inflammation due to infection, but two case reports (N Engl J Med 1992;323:1681-4; Thorax 1993;48:70-1) have suggested that CD alone (patients have absent MCC but motile cilia and normal ciliary ultrastructure) can be inherited and cause Kartagener’s syndrome (KS; primary ciliary dyskinesia). We have investigated a family in which two siblings have KS but motile cilia with normal ultrastructure. We have also treated (antibiotics and topical corticosteroids) one patient with chronic mucopurulent sinusitis and two patients with KS and CD alone to determine if CD is reversible. The mother, father and a healthy sibling did not have CD (SD of angles <15.6°), whereas the two affected siblings had CD which was unchanged in biopsies taken 10 years apart (22.6° and 26.7°; 26.4° and 25.0°). Right and left bronchial cilia from one sibling were also disoriented (23.5° and 22.2°). In sinusitis CD was reversible (21.5° to 10.3°) but in two KS patients with CD alone it was not (25.2° to 24.5°; 23.8° to 23.1°). The presence of CD in two siblings, in biopsies taken 10 years apart, in cilia from different parts of the respiratory tract, and the lack of reversibility in such patients compared with mucopurulent sinusitis, supports the hypothesis that CD alone (primary CD) can cause KS.

Ciliary disorder alone can cause primary ciliary dyskinesia syndrome

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Two case reports have proposed that ciliary disorder alone may cause the syndrome of primary ciliary dyskinesia (PCD). These cases had motile cilia and normal ciliary ultrastructure but inefficient transport because the direction of the ciliary beat was disoriented. We have identified 11 patients with the clinical features of PCD who have normal ciliary ultrastructure. A clinical assessment, chest radiograph, pulmonary function tests, nasal mucociliary clearance (NMCC), ciliary beat frequency (CBF), ciliary ultrastructure and orientation were assessed in each subject and eight of the 11 had computed tomography of the thorax. The clinical features were the same as those previously reported in patients with PCD: chronic cough and spum production, bronchiectasis, chronic sinusitis, middle ear problems, dextrocardia (in five) and infertility (one male had immotile sperm). Cilia ultrastructure was normal but NMCC was absent in all cases. The CBF ranged from 8.4 to 14.9 Hz. Ciliary beat pattern was stiff in seven cases, six of whom had slow (<11 Hz) CBF. Ciliary disorientation was measured using an Imagination processing system. The cilia were significantly disoriented in all cases when measured by both the central pair (range 21.8-26.4°) and basal feet (range 20.6-28.9°) compared with 16 normal controls (range 11.0-15.5 and 12.3-17.6°, respectively). This study suggests that ciliary disorientation alone can lead to the clinical syndrome of PCD. Ciliary disorientation should be assessed if NMCC is absent, CBF is normal or near normal, and ciliary ultrastructure is normal.

Ciliary function tests: can we detect recurrent microaspiration in children with gastro-oesophageal reflux?

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Respiratory tract disease associated with gastro-
osseophageal reflux (GOR) may be reflex vagally mediated or result from aspiration. Our aim was to determine if ciliary motility is impaired by in vitro aspiration of gastric juice (GJ) or oropharyngeal contents and hence to explore the potential of ciliary function tests in the diagnosis of recurrent microaspiration in children with GOR. Cilia from the inferior turbinates of healthy volunteers were obtained by brushing. Control ciliated epithelium was incubated in growth medium at 37°C. The remaining ciliated cells were exposed transiently (in vitro aspiration) at 37°C to milk, saliva and four solutions of GJ in which the concentration of HCl and peptic varied, followed by “rescue” into growth medium. Ciliary beat frequency (CBF) was measured using a computerised photometric technique. The odds of ciliated cellular death was calculated. Interaction between GJ components was assessed by factorial design analysis. In vitro aspiration of milk and saliva had no effect on ciliary function whereas transient exposure to GJ impaired CBF (p<0-05) and increased the odds of ciliated cellular death by 2-5–8-5 times (95% confidence interval). Increasing the HCl or peptic concentration further enhanced the ciliotoxic effect of GJ but there is no synergism. Transient exposure of ciliated epithelium to GJ significantly impairs ciliary motility. An assessment of tracheal ciliary function may be useful in distinguishing recurrent microaspiration of GJ in children with GOR from aspiration direct from the oropharynx or vagally mediated respiratory symptoms.

Breaking the “bad news” of lung cancer: a study of patients’ views and support needs

M CAMPBELL, A HUGHES, J MACFARLANE Respiratory Medicine, City Hospital, Nottingham We studied 20 consecutive patients who had been told their diagnosis of lung cancer in our combined lung oncology clinic (CLOC) within the previous two weeks and had seen our Macmillan support sister (MCS) at the same time. Our normal practice is to review the patients again in the clinic 2–3 weeks later and to telephone the GP from the clinic when bad news is broken. All patients were contacted by phone by MC within two weeks after the diagnosis discussion and asked questions in a semi-structured manner. Ten patients had “felt worse” since knowing the diagnosis, in only two cases due to deteriorating physical symptoms. Ten felt unchanged or better. Eleven patients had had no professional contact following the clinic visit, six had been contacted by their GP, and three by other staff. Eighteen felt they had understood all or most of the information given in the CLOC but 15 would have liked further discussions shortly after about the diagnosis and their concerns. Of those five patients who did not wish further discussions, three had families who did want more information. Concerns raised mainly involved fear of the treatment and investigations (eight) and uncertainty about their future (six); only two did not understand the information provided initially. Fourteen patients said they would have benefited from a home visit mostly within the first week, two would have liked a visit in the future, two were happy with the planned clinic support, and two were quite content with the support from the GP/other staff. We conclude that, although most patients understand the information given to them at the time of diagnosis, there is a big need for rapid follow up and contact from the lung cancer team, possibly in the form of a home visit by a Macmillan support sister.

Lung cancer in ex-smokers

M MUNAVAR, BR O’DRISCOLL Salford Royal Hospitals NHS Trust, Salford It is frequently stated that ex-smokers have a negligible risk of developing lung cancer once they have stopped smoking for more than 10 years (BMJ 1994;308:1479). However, Doll and Hill reported that doctors who gave up smoking even 20 years previously had a threefold greater incidence of fatal lung cancer than life long non-smokers (0-19 e 0-07 deaths per 1000) (BMJ 1964;1:1399–410). We have recorded the smoking status prospectively in 872 patients referred for fiberoptic bronchoscopy. Of these 274 (31%) were found to have a visible tumour in the central airways. The smoking status of these patients is shown in the table and the data of Doll and Hill are shown for comparison. We conclude that longstanding ex-smokers have a reduced (but not negligible) risk of lung cancer.

Risk of lung cancer following radiotherapy for breast cancer in Yorkshire

HES HOSKER, O JOHNSON, T PAYNE, C JOHNSTON, SB PEARSON Leeds Chest Clinic and Cancer Registry, Cookridge Hospital, Leeds Recent studies suggest that radiotherapy for breast cancer increases the risk of lung cancer, but there are no available data from the UK. A retrospective case note study was set up to examine any excess risk in patients with breast cancer treated with radiotherapy in Yorkshire. All patients with proven breast cancer between 1975 and 1984 and a subsequent diagnosis of lung cancer at least six months later were identified. Patients with inadequate data, or lung histology suggesting adenocarcinoma, were excluded. Patients were compared with 660 matched case controls with breast cancer diagnosed in the same year who had not developed lung cancer. The number of patients from each group who had received radiotherapy for breast cancer was compared. Ninety nine cases were identified with breast cancer and subsequent lung cancer. Thirty three of these had proven lung cancer histology which was not adenocarcinoma, 21 (64%) of whom had received radiotherapy for breast cancer, compared with 288 of 660 (44%) of the case controls who had not developed lung cancer. The relative risk of prior radiotherapy was 2-3 (95% CI 1-1 to 4-9; p=0.03). Histological types of lung cancer were similar in the radiotherapy and non-radiotherapy groups, with 61% being squamous, 27% small cell, and 9% poorly differentiated. There was one case of non-Hodgkin’s lymphoma. These patients with breast cancer who subsequently develop lung cancer are twice as likely to have received radiotherapy for breast cancer than patients who do not develop lung cancer, supporting the hypothesis that such radiotherapy increases the risk of subsequent lung cancer.

Completion pneumonectomy for lung cancer: indications and outcome

K AL KATTAN, P GOLDSTRAW Royal Brompton Hospital, London Completion pneumonectomy for lung cancer has been associated with higher rates of mortality and morbidity and this is reflected in the selection of cases. Over a period of 14 years (January 1980 to December 1993) 24 completion pneumonectomies for lung cancer were performed representing 4-5% of all pneumonectomies for lung cancer. There were 15 right and nine left completion pneumonectomies (CP) for an average age of 64 years (range 39–77). The indication for CP included local recurrence in 10, a second in nine, and a new primary after previous pulmonary resection for tuberculosis in two patients. Operative complications requiring CP after resection for lung cancer included two bronchial strictures and one empyema. There was one operative death and 6-7% morbidity in form of bleeding in two, prolonged ventilation in one and empyema in one. We conclude that CP can be performed with an acceptable risk in selected patients. This aggressive approach should only be performed as a potentially curative procedure.

Value of cervical mediastinoscopy combined with anterior mediastinotomy in bronchogenic carcinoma of the left upper lobe

X JIAO, P GOLDSTRAW Royal Brompton Hospital, London In the pre-operative evaluation of a patient with lung cancer assessment of the mediastinum to exclude invasion and proximal nodal disease remains a fundamental importance. In this report, cervical mediastinoscopy is of proven benefit but the development of scanning techniques have allowed this invasive procedure to be applied selectively. The situation is complicated for patients with tumours within the left upper lobe since cervical mediastinoscopy cannot exclude nodal disease beneath the aortic arch, lateral to the aortic arch or within the anterior mediastinum, nor exclude direct tumour invasion in this area. The addition of left anterior mediastinotomy allows biopsy of the subaortic lymph nodes, bidirectional palpation of the subaortic fossa and, if necessary, the intrapericardial exploration. We have sought to evaluate...
the contribution that this additional minor operation makes to the preoperative selection of patients with tumours within the left upper lobe. From January 1990 to July 1994, 85 patients who were otherwise thought to have an operable tumour within the left upper lobe underwent cervical mediastinoscopy and left anterior mediastinotomy. Twenty seven (31%-8%) patients were found to be inoperable, either because of nodal involvement at cervical mediastinoscopy (four patients) or because of extension into the mediastinum at left anterior mediastinotomy (14 patients), or because of positive results from both methods (nine patients). The inoperability determined by this examination for patients with adenocarcinoma (8/18, 44%-4%) is higher than for patients with squamous carcinoma (12/25, 31%-3%). Of the 58 patients with negative findings all proceeded to thoracotomy and complete resection was possible in 54 patients (93%-3%). We conclude that left anterior mediastinotomy is a valuable additional staging investigation for patients with tumours originating within the left upper lobe.

Thymoma: a comparison between surgical and pathological classification

K A L KATTAN, M SWEETS, M SHEPARD, P GOLDSWORTH Royal Brompton Hospital, London During the period of 13 years between January 1981 and December 1993 54 resections were performed for thymic tumours. A complete revision of the histological slides was available for 27 patients with true thymoma. Intraoperative clinical staging showed 14 in stage 1, three in stage 2, six in stage 3, and four in stage 4. Average follow up period of 63 months (24-109) showed significant longer survival in clinical stage 1 than stages 3 and 4 (actuarial 10 year survival 91% compared with 66% and 50%, respectively). The Rosai-Levine cellular staging and the Muller classification failed to correlate with long term survival compared with the Masaoka and operative clinical staging. Complete surgical resection showed a better survival. We conclude that the Masaoka and clinical staging is the best prognostic indicator; however, multicentre studies are needed to evaluate the other classifications. For true thymoma complete surgical resection should be performed to ensure long term survival.

Pleural lactic dehydrogenase (LD) and isoenzymes as additional diagnostic markers in pleural effusion

D DEV, J JOSEPH, MJ SMITH, SM VINEY, P BECK, GS BASLAN Respiratory Unit, South Yorkshire Department of Biochemistry, Rotherham General Hospital NHS Trust, Rotherham, South Yorkshire Pleural fluid total LD analysis is being increasingly used in the USA and Europe in separating transudative from exudative pleural effusions. The further role of LD and its isoenzymes as an additional marker to aid in the differential diagnosis of pleural effusions has not been extensively explored. We have studied the biochemical profile of 100 patients with pleural effusions due to a variety of defined diagnoses with special reference to total LD (IU/L), LD isoenzymes, and total protein (TP). Total LD, LD fluid/serum (F/S) ratio and LD isoenzymes (1-5) F/S ratio were found to be minimum in cardiac failure (CCF) and maximum in empyema. The value was intermediate in malignancy and other exudative conditions as shown in the table (median concentrations).

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>TP (IU/L)</th>
<th>F/S ratio</th>
<th>Total LD (IU/L)</th>
<th>F/S ratio</th>
<th>LD-5 F/S ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cardiac failure</td>
<td>37</td>
<td>0-37</td>
<td>95</td>
<td>0-4</td>
<td>0-79</td>
<td></td>
</tr>
<tr>
<td>Empyema</td>
<td>17</td>
<td>0-70</td>
<td>3950</td>
<td>10-17</td>
<td>18-9</td>
<td></td>
</tr>
<tr>
<td>NMT</td>
<td>65</td>
<td>0-61</td>
<td>281</td>
<td>1-1</td>
<td>1-4</td>
<td></td>
</tr>
<tr>
<td>Mesothelioma</td>
<td>7</td>
<td>0-59</td>
<td>157</td>
<td>0-89</td>
<td>2-8</td>
<td></td>
</tr>
<tr>
<td>Pulmonary effusion</td>
<td>20</td>
<td>0-60</td>
<td>100</td>
<td>0-97</td>
<td>1-2</td>
<td></td>
</tr>
<tr>
<td>Parapneumonic</td>
<td>12</td>
<td>0-52</td>
<td>274</td>
<td>0-9</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Others</td>
<td>22</td>
<td>0-64</td>
<td>469</td>
<td>1-65</td>
<td>4-2</td>
<td></td>
</tr>
</tbody>
</table>

The difference between CCF and non-CCF was significant (p<0.001). Isoenzyme LD-5 ratio tended to be higher (p<0.05) in pleural effusions of mesothelioma origin compared those from non-malignant pleural effusions (NMT). Thus, for patients with cardiac failure presenting with pleural effusion, the finding of a total LD <100 IU/L combined with a TP F/S ratio of <0.4 excluded other serious clinical conditions such as pulmonary embolism, parapneumonia, and malignancy as a cause of the effusion.

Differential pleural effusion kinetics in mesothelioma and carcinoma: a study using a dual isotope technique

D DEV, J JOSEPH, MJ SMITH, GS BASLAN Respiratory Unit, Rotherham General Hospital, Rotherham, South Yorkshire Pleural fluid effusion results from an imbalance between the rate of formation and rate of removal of fluid from the pleural space. We have monitored the formation and drainage of pleural fluid in man using a radiolabelled protein technique (Nucl Med Commun 1992;13:432). Fifty three patients with free flowing pleural effusion from a variety of causes were investigated using this method. Fluid formation was monitored by measuring the rate of movement from blood into effusion of transferrin radiolabelled in vivo with iodine-125 [Kout]. Fluid drainage [Kout] was estimated by monitoring the rate of appearance in the blood of radiolabelled albumin (iodine-125) injected directly into the effusion. The Kout and Kout were calculated using a three compartmental model and expressed in the table as mean (SE) (10^-4 h^-1). Outflow (Kout) was significantly lower in effusions due to mesothelioma than primary lung cancer (p<0.005), metastatic lung cancer (p<0.001), or cardiac failure (p<0.001). The inflow (Kinf) was maximum in pancreatitis (Kin of 114) consistent with increased vascular permeability/exudation. We speculate that, in mesothelioma, the predominant mechanism of effusion formation is the effect of the tumour on the lymphatic system, resulting in low Kout, even before the tumour produces the gross pleural encasement. In carcinomatous involvement of the pleura, however, the predominant mechanism appears to be a modest increase in rate of exudation due to increased vascular permeability. Because of the apparent difference in mechanism and consequently lower values of Kout, the isotope technique could form the basis of a simple methodology for distinguishing malignant pleural effusions associated with mesothelioma from those associated with carcinoma— a common clinical dilemma.

Use of genetic markers to identify precancerous bronchial lesions in lung carcinogenesis

U PASTORINO, P GOLDSWORTH Royal Brompton Hospital, London In an attempt to define premalignant changes occurring in the multistep process of lung carcinogenesis, a cytogenetic and genetic study was performed at the National Cancer Institute of Milan on 94 patients undergoing pulmonary resection for an early stage lung cancer and 11 controls resected for other diseases. The panel of genetic markers included the assessment of chromosomal abnormalities, overexpression of EGFR and HER2/NEU and p53 mutation in normal bronchial epithelium and primary tumour specimens. Of 94 cases, 40 displayed multiple synchronous or metachronous tumours in the upper aerodigestive field, while 54 had either single tumours (45) or multiple tumours out of field (9). Cytogenetic alterations were observed in 60% (28/47) of the evaluable tumour specimens with common rearranged karyotypes, particularly involving chromosomes 3 (64%) and 17 (36%). Gene alterations were detected including overexpression of the EGFR in 57% (46/80), HER2/NEU in 21% (17/79), and p53 mutations in 47% (23/49). The overall frequency of genetic changes (any type) in the tumour was 78% (73/94). In the normal bronchial mucosa we identified a rearranged karyotype in 23% of the evaluable cases (20/86); particularly simple rearrangements involving chromosomes 3 (13 cases), 7 (six cases), and 17 (three cases), as well as overexpression of EGFR in 35% (27/74) and of HER2/NEU in 15% (11/74). The overall frequency of genetic changes (any type) in the normal epithelium was 43% (39/91). The overall frequency of genetic changes in the normal epithelium was higher in patients with multiple tumours in the field (60%) compared with those with single or multiple tumours in other sites (30%). In addition, deletions of chromosome 3p and p53 mutations have been detected in precancerous lesions of the bronchial mucosa in five patients. These data indicate that various stages of lung tumorigenesis display specific genetic alterations and that a "field carcinogenesis" effect results in
the appearance of detectable genetic lesions at sites distant from the tumour.

Acute neurohormonal responses to hypoxaemia in humans

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Whilst the acute pulmonary pressor response to hypoxaemia in man is well documented, there remains some controversy regarding the effects of this stimulus on integrated hormonal responses, particularly on activity of the renin-angiotensin-aldosterone system (RAAS). This may be important in the adaptive response to alveolar hypoxia, either in pulmonary disease or adverse environmental conditions. These neurohormonal changes have not been fully characterised nor studied in association with the haemodynamic effects of hypoxaemia. These responses were therefore studied in 10 healthy volunteers on two separate occasions. After reaching a resting haemodynamic state, subjects breathed for 30 minutes either room air or a nitrogen/oxygen mixture which rendered SaO2 between 75% and 80%. Pulmonary and systemic haemodynamic parameters were measured and venous blood samples taken at baseline and after 30 minutes breathing air or the hypoxic gas. Hypoxaemia significantly altered heart rate, cardiac output, and mean pulmonary artery pressure (MPAP), but not mean arterial pressure (MAP) compared with normoxaemia. Although plasma renin activity (PRA) and angiotensin II were unaffected by hypoxaemia, plasma aldosterone (Aldo) fell significantly in comparison with normoxaemia. There was a non-significant increase in plasma levels of cortisol and catecholamines during hypoxaemia but no changes were observed during normoxaemia (table). Thus, the renin-angiotensin system becomes dissociated during hypoxaemia where plasma aldosterone levels decreased despite having no significant effects on other components of the RAAS or other markers of adrenal cortical and medullary function. This dissociation may be influenced by the counter-regulatory effects of the natriuretic peptides released under hypoxic conditions.

Effect of acute changes in oxygen tension on responses to methacholine in bronchi from chronically hypoxic rats

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Sustained alveolar hypoxia is a feature of airway diseases such as chronic obstructive pulmonary disease. We have previously shown in bovine bronchi (Clayton et al. Br J Pharmacol 1994;112:593P) that acute changes in O2 levels alter reactivity to methacholine (MCh); however, it remains to be seen whether chronic hypoxia alters the sensitivity of bronchi to acute changes in O2 tension in the rat. We examined contractions of chronically hypoxic and control rat bronchi to MCh at 95%, 20%, and 4% O2. Rats were reared in a hypoxic/hypobaric chamber for 14 days under a pressure of 500–550 mmHg, giving effective gas concentrations of 10% O2, 0·3% CO2, and balance N2. Rats were sacrificed and contractions of rings of first order bronchi were measured isometrically (n = 6 in each case). Cumulative concentration–response curves were constructed to MCh (10−6–3 × 10−4 M), results being expressed as % of the maximum response in 95% O2. In control rats there was no difference between responses in 95% and 20% O2, however, responses in 4% were significantly (p<0·001) attenuated, with a mean 29·3% decrease in maximum response. Bronchi from hypoxic rats showed a similar pattern; with responses in 4% being significantly (p<0·05) attenuated with a mean 25·6% decrease in maximum response. Exposure to chronic hypoxia therefore does not appear to alter sensitivity to acute changes in O2 tension in rat bronchi. [Supported by the Chest Heart and Stoke Association (Scotland), the National Foundation, and the National Asthma Campaign.]

Effect of hypoxia and β agonists on the activity of the renin-angiotensin system

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We have reported that the renin-angiotensin system (RAS) is activated in acute severe asthma and also by β agonist therapy on plasma renin and angiotensin II (AII) levels. On four separate study days in a double blind, placebo controlled crossover study eight healthy volunteers were randomised to receive either hypoxic mixture (FiO2 12%) or air for a period of 30 minutes, with either salbutamol (5 mg) or placebo (3·5 ml N salmine) administered into the circuit via a nebuliser (at time t = 10 min). Oxygen saturation was monitored throughout the study by pulse oximeter and blood withdrawal at baseline, 10, 20, 30, 45, 60, 90, and 120 minutes for measurement of plasma renin and AII. There was no significant difference between baseline values of renin or AII on either study day. Following the period of hypoxia alone [mean (SE) O2 saturation 82·8 (1·5), 87·6 (3·9), 84·5 (3·0)% at 10, 20 and 30 minutes] there were no significant changes in plasma levels of renin or AII. When salbutamol was added to the hypoxic mixture, there were significant rises in plasma renin and AII levels [mean (SE) increase in AII from baseline of 7·2 (9·9), 4·5 (3·4), 5·4 (2·9), 3·6 (3·1) pmol/ml at 30, 45, 60, and 90 minutes, and in renin of 141 (63), 128 (4), 155 (63), 40 (3·2) and 40 (3·9) pmol/ml at 30, 45, 60, 90, and 120 minutes, respectively]. Nebulised salbutamol in air also increased plasma renin and AII levels, but there was no significant difference between the effect of salbutamol in either the normoxic or hypoxic mixture. We conclude that there is activation of the renin-angiotensin system by hypoxia, but not by hypoxia. Hypoxia does not influence the effect of salbutamol on the RAS. Thus, β agonists, but not hypoxia, are likely to be contributing to the activation of the RAS in acute asthma. [Supported by the National Asthma Campaign.]

Angiotensin II (AII) potentiates, in vitro and in vivo, methacholine (MCh)-induced bronchoconstriction

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Plasma AII levels are raised in patients with acute severe asthma and also by β agonists. In addition, intravenous AII evokes bronchoconstriction in mild asthmatic patients. This study examined the effects of AII on bronchi in vitro and the interaction of AII with MCh-evoked contractions both in vitro and in vivo. Contractions of rings of human and bovine bronchi were measured isometrically. Cumulative concentration–response curves were obtained to AII (10−8–10−4 M). In addition, seven asthmatic patients with mild bronchial hyperreactivity to MCh were studied. In a randomised double blind study, patients received placebo or AII 1 or 2 mg/kg/min by intravenous infusion. These doses were below those which evoke bronchoconstriction. At the end of the infusion MCh challenge was undertaken. FEV1 values were measured at baseline, at the end of the infusion, and following each MCh dose. PC20 responses were expressed as geometric mean and range. AII alone in vitro evoked only small (<25 g) contractions of human and bovine bronchi (threshold for contraction, 3 × 10−5–3 × 10−7 M). Pre-incubation with AII 10−4 M significantly (p<0·001) enhanced contractions to MCh with an increased maximum of 39.1% in human and 13.7% in bovine bronchi (n = 6 in each case). In mild asthmatic patients AII alone evoked no change in baseline FEV1 values. The geometric mean PC20 of MCh after placebo infusion was 3·09 mg/ml (range 1.15–6·0 mg/ml). After infusion with AII 1 mg/kg/min this was 2·14 mg/ml (range 0.85–3·8 mg/ml). In the presence of AII 2 mg/kg/min there was a significant (p = 0·006) decrease in PC20 compared with placebo (geometric mean, 1·2 mg/ml, range 0·45–2·08 mg/ml). In conclusion, AII in susceptible patients evoked bronchoconstrictions both in vitro in human and bovine tissue, and also in vivo in mild asthmatic patients. [This work was supported by the National Asthma Campaign.]

Lisinopril inhibits elevation of plasma angiotensin II by nebulised β agonists

EA MILLAR, GT MCINNES, NC THOMSON Department of Respiratory Medicine, Western Infirmary, Glasgow

We have described activation...
endopeptidase-24-11 (NEP) and binding to a non-guanine cyclase clearance receptor. We have found in studies on human airway that phosphoramidon, whose actions include inhibition of NEP, enhances the in vitro protective effect of ANP on methacholine induced contraction. Similarly in vivo thoriphan, an inhibitor of NEP, enhances the protective effect of ANP against histamine-induced bronchostenosis. We therefore hypothesised that pretreatment with thoriphan might enhance the bronchodilator response to inhaled ANP. In a randomised, double blind, placebo controlled, crossover study, six asthmatic patients (one woman), mean (SD) age 47-3 (3-8) years and FEV1 91-0 (42-2), 55 (3-8) % predicted, were studied. All were shown at screening to have at least a 25% improvement in FEV1 to salbutamol. Patients were then randomised to receive for six hours each of placebo, dexamethasone, or two doses of salbutamol. On study visits they received either thoriphan 1 mg (in 2 ml) followed by ANP 5 mg or placebo (saline) or placebo (saline) followed by ANP (5 mg), placebo or salbutamol 5 mg. Spirometric values were measured after each inhalation then followed for two hours. ANP alone caused a significant bronchodilator response at 10 and 15 minutes when compared with placebo and thoriphan alone, mean (SE) % change in FEV1, of 16-8 (8-1) and 16-1 (6-8), respectively. Prior inhalation of thoriphan prolonged bronchodilator effect of ANP, significant bronchodilatation being noted at five minutes (mean (SE) % change in FEV1 of 17-5 (1-2)) and being maintained up to 60 minutes (23-1 (3-4)). With both ANP and the combination of the NEP inhibitor and ANP the degree of bronchodilation was significantly less than that produced by salbutamol which was maximal at one hour (mean (SE) % change in FEV1 of 53 (10-5)) and maintained at two hours. These results confirm that airway NEP is important in modulating the effect of inhaled ANP. [Supported by UCB and the British Lung Foundation.]

Bromodilator responsiveness to salbutamol and histamine reactivity after continuous treatment with twice daily salmeterol

A GROVE, RA CLARE, JH WINTER, DP DHILLON, LC MCFARLANE, D MCDEVITT, BJ LIPWORTH Department of Clinical Pharmacology and Respiratory Medicine, University of Dundee Medical School, Ninewells Hospital, Dundee Seventeen asthmatics, all receiving inhaled steroids, of mean (SD) age 34 (3) years and FEV1, 62 (3%) % predicted, were evaluated. After a two week run in without β2 agonists they received salmeterol (SMT) dry powder 50 μg bd or placebo (PL) for four weeks in a randomised double blind crossover design. Twelve hours after the last dose of each treatment a histamine test was performed and after 36 hours a salbutamol DRC was constructed. Airway and systemic baselines were not significantly different for SMT v PL. There was a partial right shift in DRC after SMT v PL for PEFR, p<0.036 and FEV1, p<0.058, but not for FEF25-75. The maximum bronchodilator delta response was not, however, attenuated: FEV1, SMT 0.901 PL 0.701 (95% CI 0-2 to 0-24), PEFR, SMT 100/l/min PL 116/l/min (95% CI 6 to 38/min). The FEV1, PEFR (log) following SMT was 0.37 mg/ml v PL 0.20 mg/ml, p<0.05 (95% CI 0-002 to 0-334). Systemic responses were attenuated after SMT PL (p<0.05): potassium: SMT 0.73 mmol/l PL 1.03 mmol/l (95% CI 0-05 to 0-05 mmol/l). Lymphocyte β2 density (log) tended to be lower after SMT 0.22 fmol/106 cells compared with PL 0.31 fmol/106 cells, or after run in 0.31 fmol/106 cells (NS) and PEFR was improved during treatment with SMT v PL (p<0.05): 421/l/min v 395/l/min (95% CI 6 to 46/min). In summary, SMT 50 μg bd produced a right shift in salbutamol DRC but did not blunt the maximal bronchodilator response, resulted in a small but significant protection against histamine reactivity, significantly blunted the maximal systemic β2 responses to salbutamol, and showed a trend to reduced lymphocyte β2 adrenoceptor density.

Effect of cyclosporin A (CaA) on the expression of activation markers by T lymphocytes in chronic severe asthma

SH LOCK, CJ CORRIGAN, NC BARNES, AB KAY Royal Brompton National Heart and Lung Institute, Department of Allergy and Clinical Immunology, London, The London Chest Hospital, and the University of Warwick. We have shown that elevated numbers of peripheral blood T lymphocytes express activation markers in acute severe asthma and the reduction of expression after therapy could be correlated with improvement in lung function (Am Rev Respir Dis 1990;141:970–7). We have studied the expression of activation markers by peripheral blood T lymphocytes in...
in 39 corticosteroid dependent asthmatics taking part in a double blind, placebo controlled corticosteroid reduction study using CsA. Nineteen identical placebo. Proceeding of in significant medication there obstructive lymphocytes were measured using flow cytometry. There was no significant difference in activation marker expression between the cyclosporin and placebo groups at baseline. After 24 weeks of trial medication there was a significant reduction in HLA-DR (median 13%, \(p=0.01\)) and CD25 (\(-37\%, p<0.01\)) expression on CD4 T lymphocytes in the CsA group compared with the placebo group. CsA reduces the expression of activation markers on peripheral blood T lymphocytes from asthmatics and this may be central to the mechanism of action of CsA in asthma.

MRI cerebral white matter changes in obstructive sleep apnoea (OSA)

RJO DAVIES, S RENOWNDEN, N MOORE, JR STRADLING Osher Chest Unit and Radiology Department, Oxford Radcliffe Hospital, Oxford Patients with obstructive sleep apnoea suffer an excess of cerebrovascular disease, but this might be due to many confounding risk factors seen in these patients, particularly obesity. Cerebral magnetic resonance imaging (MRI) allows the identification of clinically silent cerebral ischaemic demyelination which is seen as small areas of white matter high signal. To assess whether such silent ischaemia is more frequent in OSA patients than in controls matched for the major confounding variables, eight patients with OSA and 10 control subjects underwent MRI. The study groups did not differ for any of the following indices: sex, age, body mass index, alcohol and tobacco intake, daytime ambulatory blood pressure, waist to hip ratio, and fasting lipid concentrations. The patients with OSA had their obstructive sleep apnoea confirmed by full polysomnography (\(4\%\) SaO\(_2\), dip rate \(>15\)/hour). The controls were normal community subjects. All subjects underwent T1 and T2 weighted MRI imaging. These were acquired 39-72 hours after performance and interpreted blind of the patient/control status. The presence of white matter high signal abnormalities was scored at three levels: (1) no abnormality, (2) 1-5 abnormal foci, (3) \(>5\) abnormal foci (table).

<table>
<thead>
<tr>
<th>Number of abnormal high signal foci</th>
<th>None (1) (5) (&gt;5)</th>
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<tr>
<td>OSA patients</td>
<td>5</td>
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<tr>
<td>Controls</td>
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</table>

Both the patients with OSA and the matched control subjects showed frequent cerebral ischaemic lesions consistent with their generally unhealthy state. These were no more prevalent in the patient group than the control subjects.

Nasal CPAP as a treatment for obstructive sleep apnoea: objective versus subjective compliance

JMPARTLETT, DJ PITS ON, S BUCK, JR STRADLING Osher Chest Unit, Churchill Hospital, Oxford Nasal continuous positive airway pressure (CPAP) successfully treats obstructive sleep apnoea (OSA). Data from the US (Kribbs et al. Am Rev Respir Dis 1993;147:887) has suggested poor compliance measured objectively under 25 hours/week. It is not clear whether compliance needs to be measured or whether patients report usage accurately. This study was carried out on 71 patients using CPAP over an average period of 25 months (range 9-60). The patients were unaware that the clocks were being read for the study. Subjective compliance was collected as part of a validated questionnaire survey (Hoffstein et al. Am Rev Respir Dis 1992;145:841) which allowed comparison with this Canadian group. Mean (SD) objective compliance in our patients was 41.2 (11.7) hours/week (5-9 hours/night), and subjective compliance was 49.6 (9) hours/week (7-9 hours/night). The clock was worn for 18 hours in the study. On average, our patients overreported their usage by 8-35 (95% CI 5-7 to 11-0) hours/week. The graph illustrates the relationship between subjective and objective compliance. A subgroup \((n=21)\) had two timerock readings one year apart; mean

Variation in CPAP requirement using an intelligent CPAP device

N POTTER, AK SIMMONDS Royal Brompton Hospital, London Although conventional continuous positive airway pressure (CPAP) machines deliver a constant pressure throughout the night, the pressure required to maintain airway patency in any individual with obstructive sleep apnoea (OSA) may vary depending on a number of factors including sleep stage and body posture. The Sullivan Autoset Clinical system (Rescare) has been developed to titrate CPAP level to nasal airflow. It has been used in this pilot study to assess the individual variation in overnight CPAP requirement in 17 patients (15 men) with baseline apnoea/hypopnoea index (AHI) of 33 (range 7-91). In eight patients sleep staging was carried out during the autoset study and body position monitored by video recording. For the group as a whole median (SD) CPAP requirement was 8.1 (1.2) cmH\(_2\)O with a 90th centile range of 10.9 (1.6) cmH\(_2\)O. Mean (SD) apnoea index determined by the autoset was 4 (4.8). There was, however, a wide range in CPAP requirement within individuals overnight. In some patients CPAP level was related to sleep stage, rising in REM sleep, in others changes in sleep stage had no consistent effect. Two groups were identified: one with a \(<2\) cmH\(_2\)O variation between median and 90th centile CPAP level and the other with a \(>2\) cmH\(_2\)O variation between these values. Body mass index did not differ significantly between the groups but the group with the greater variation in overnight CPAP requirement had a higher baseline AHI (50 vs 39, \(p=0.01\)) indicating more severe disease. The consequences of failing to accurately titrate CPAP are unknown, but there does seem to be a subgroup of OSA patients with a highly variable CPAP requirement which might benefit from more flexible overnight treatment.

Use of the Epworth Sleepiness Scale (ESS) to monitor response to treatment with nasal continuous positive airway pressure (nasal CPAP) in patients with obstructive sleep apnoea

FM HARDINDE, D PITS ON, JR STRADLING Osher Chest Unit, Churchill Hospital, Oxford The ESS (Johns. Sleep 1991;14:50-5) is a questionnaire derived scale of sleepiness which can be used to assess the degree of daytime sleepiness suffered by patients with obstructive sleep apnoea (OSA). Its use has been well validated in a variety of sleep conditions. When patients with OSA are treated with nasal CPAP their subjective sleepiness often markedly improves. But objective tests of daytime vigilance such as the multiple unprepared reaction time test or multiple sleep latency test have demonstrated significant but often only very small improvements following treatment with nasal CPAP. We administered ESS questionnaires to patients at initial assessment and at two months after commencing nasal CPAP in one group (48 patients, mean age 50-2 years, 94% men), and after one year of treatment with nasal CPAP in a second group (18 patients, mean age 49-7 years, all men). In the first group mean (SE) ESS before treatment was 16 (0-5) and after two months treatment had fallen to 8 (0-6) (\(p=0.0001\)). In the second group mean ESS before treatment was 14 (1-3) and after treatment for one year had fallen to 6 (0-9) (\(p=0.0001\)). In this second group compliance with treatment was monitored in 12 patients using Respironics machines with time clocks of which patients were unaware. Mean (SE) compliance was 6:54 (0-52) hours/night. We conclude that changes in the ESS following treatment with nasal CPAP are more representative of the magnitude of patients' subjective response to treatment than other
objective tests of daytime vigilance, possibly because the ESS is measuring a different component of sleepiness. It is therefore likely that the ESS could be useful clinically to monitor progress during treatment.

Use of pulse transit time as a measure of inspiratory effort in obstructive sleep apnoea

DJ PITSON, A SANDELL, R VAN DEN HOUT, JR STRADLING Oiler Chest Unit, Churchill Hospital, Oxford Pulse transit time (PTT) is the time taken for the arterial pulse shock wave to travel from the aortic valve to the periphery. PTT is inversely proportional to blood pressure and it has been shown that, like systolic blood pressure swings, the size of the inspiratory swings in PTT correlate well with the degree of inspiratory effort in awake normal subjects breathing through an added threshold inspiratory valve. This study investigated the ability of inspiratory PTT swings to act as a measure of inspiratory effort in patients with obstructive sleep apnoea (OSA). Recordings were made on eight men of mean age 44 (range 34–58) years with OSA (mean >4% SaO2 dip rate 43, range 13–68 dips/hour) during their initial titration of nasal continuous positive airway pressure (NCPAP). NCPAP was decreased from the optimal criterion used. A study was performed to determine pressure at three minutes intervals in steps of 1 cm H2O to produce a range of inspiratory efforts. Oesophageal pressure (IOP) and PTT were included in the standard sleep monitoring. About 20 breaths at each NCPAP level were measured for each patient. IOP swings were grouped together in sampling bins of 5 cm H2O (range 0 to −65 cm H2O) and the mean and standard error of the PTT swings within each bin was calculated. The slopes, intercepts and r values are shown in the table. The correlation between PTT swings and analysed, oximetry at outpatients followed by polysomnography offers a lower financial cost than generalised polysomnography. In a selected population method B is preferable; in a non-selected population method C is the best option.

Comparison of laboratory polysomnography with home use of the Edentrace II recorder in the diagnosis of sleep apnoea

SP FINCH, IL MORTMORE, NL DOUGLAS Sleep Laboratory, Royal Infirmary, Edinburgh Great awareness of sleep apnoea, both within the medical profession and the public, has resulted in a steady increase in the number of referrals to sleep laboratories. Home sleep monitoring is an obvious method of increasing patient assessment and thus reducing waiting lists. We have, therefore, compared full polysomnography in our laboratory with home monitoring using the Edentrace II system (Edentec Corporation) which monitors airflow, heart rate, chest wall movement and O2 saturation. Fourteen patients had laboratory polysomnography and were instructed in the use of the Edentrace system for home monitoring. Staff, patient and equipment costs, together with sleep study indices, were compared. Two home studies failed due to machine faults, and there were no failures due to patient connection errors. In the 10 patients with successful home studies there was no significant difference in apnoea/hypopnoea index (AHI) per hour in bed (Edentrace mean SE 23 (4); polysomnography 21 (4); p=0.4) or between AHI per hour in bed for Edentrace and AHI per hour of sleep assessed by polysomnography (29 (6); p=0.7). Staff time was significantly longer for polysomnography (600 (19) minutes) than for home Edentrace (47 (3) minutes; p<0.001). Home Edentrace monitoring required two laboratory visits per patient, thus travel costs were twice that of polysomnography (£14.50 vs £7.25/patient). However travel costs were limited by only using patients within a 20 mile radius of the laboratory. The overall cost (staff, travel and equipment) of home Edentrace monitoring was 33% of that of polysomnography. In this ongoing study the portable Edentrace II system seems to be a cost effective method of home diagnosis for SAHS giving comparable results for full polysomnography, but no data were obtained in fourteen of 18 subjects in this ongoing study.

Nocturnal oximetry and transcutaneous CO2 in patients with cystic fibrosis

DL SMITH, DE STABLEFORD Adult Cystic Fibrosis Unit, Birmingham Heartlands Hospital, Birmingham A not well validated method of monitoring nocturnal hypoxaemia is transcutaneous CO2 (PtcCO2) monitoring. There is now a well recognised phenomenon in cystic fibrosis (CF). In 35 patients studied at the end of a course of hospital treatment and 22 patients studied at home we found significant nocturnal hypoxaemia (SNH, defined as a percentage of total study time with an SaO2 of <90% or >10% in 25% of patients. These patients may benefit from correction of nocturnal hypoxaemia with the administration of oxygen. We have studied the characteristics of nocturnal CO2 traces in 18 CF patients with moderate to severe CF (%FEV1, 16–53, mean 31). Transcutaneous CO2 tensions (PtcCO2) measurements were made with a TCM3 system (Radiometer, Copenhagen) previously validated against arterial samples in 10 patients studied overnight. Six of the 18 patients studied exhibited SNH. Mean PtcCO2 ranged from 3.75 to 7.0 kPa. Five patients had a mean PtcCO2 of 6.0 kPa or more. The maximum overnight rise in PtcCO2 ranged from 0.2 to 1.0 kPa. The pattern of overnight PtcCO2 recordings showed a basically stable trace with episodic elevations associated with simultaneous falls in SaO2. However, not all falls in SaO2 were associated with elevations in PtcCO2 suggesting that at least two different mechanisms might account for the episodes of nocturnal desaturation seen. Spirometric parameters were not correlated with any measurements of PtcCO2 however, minimum nocturnal SaO2 was correlated with maximum PtcCO2 and maximum rise in PtcCO2 (maximum PtcCO2−mean PtcCO2). Six patients with SNH were further studied during admission of nocturnal oxygen therapy at a flow rate of 2 l/min. SNH was abolished in five and greatly ameliorated in the other patient. Oxygen administration resulted in an increase in PtcCO2 in all patients ranging from 0.4 to 1.6 kPa (mean 0.8 kPa).

The role of CD10 in the modulation of neutrophil function

M MIKAMI, GC LLEWELLYN-JONES, RA STOCKLEY Lung Immuno- biochemical Research Laboratory, The General Hospital, St Helens

Subject Slope Intercept r

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IOP swings was high, therefore inspiratory PTT swings may be a useful indirect measure of inspiratory effort in patients with OSA.

Comparative study of cost of polysomnography and oximetry in the sleep apnoea syndrome

M MAYOS, J LIEZ BETORET, L HERNANDEZ, P CASAN, J SANCHIS Unitat Funció Pulmonar, Departament de Pneumologia, Hospital Sta Creu i de St Pau; Servei Català de la Salut, Barcelona, Spain In the sleep apnoea syndrome (SAS) polysomnography (psm) is the most widely used diagnostic test. Oximetry (pox) has been proposed as an alternative test which simplifies diagnosis. Its sensitivity (S) and specificity (SP) vary according to the analytical criterion used. A study was performed to compare the financial cost of diagnosing SAS in patients by means of three defined diagnostic strategies: (A) polysomnography alone; (B) oximetry interpreted with criteria of maximum specificity, plus polysomnography if negative; (C) oximetry interpreted with criteria of maximum sensitivity, plus polysomnography if positive. The financial analysis took into account the costs of installation, maintenance and replacement of material, and staff costs. Theoretical values of S, SP and prior probability of suffering from SAS (P) were used. The average cost of diagnosing SAS (C Sas) was calculated by C Sas = C Sas/P for method A; by C Sas = C Sas + [(1-SP) (1-S) P + (SP) (1-S)] C Sas/P for B; by C Sas = C Sas + [(1-P) (1-SP) P + (1-SP) (1-S)] C Sas/P for C. The results are shown in the table. We conclude that, in the combinations of S, SP and P

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<th>P</th>
<th>C Sas A</th>
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*Oximetry at outpatients. In cases of hospitalisation, B and C are greater than A.
Lan, Birmingham
CD10/NEP (neutral endopeptidase) is a zinc metalloproteinase originally identified on acute lymphoblastic leukemia cells and more recently found on neutrophil membranes where it is thought to have a modulating effect on neutrophil function (Connelly et al. Proc Natl Acad Sci USA 1985; 82:8737).
We have therefore studied the role of CD10 in neutrophil chemotaxis, degranulation and superoxide generation. In order to do this we have studied the effects of the metalloproteinase inhibitor, 1,10-phenanthroline and a monoclonal antibody to CD10. 1,10-phenanthroline reduced neutrophil chemotaxis to 10 nM FMLP from a mean of 40-7 (4-8) cells/field to 1.2 (0-2) cells/field in a dose dependent manner (p<0.002). In addition, 1,10-phenanthroline (1 nM) pre-incubated with cells resulted in a shift of the FMLP dose response for neutrophil chemotaxis to the left. However, 1,10-phenanthroline had only a slight inhibitory effect on neutrophil chemotaxis to IL-8 with no shift of the IL-8 dose response. Similarly the anti-CD10 antibody reduced the neutrophil chemotaxis to 10 nM FMLP significantly (p<0.04) in a dose dependent manner. A control solution of 26-8 (1.8) to 14-7 (1-1) cells/field at antibody concentration of 25 mg/mL and shifted the dose response to the left. The IL-8 effect was also reduced by the antibody but there was no associated shift in the dose response. Neither extracellular proteolysis nor superoxide generation was blocked by either agent. These data suggest that the metalloproteinase, CD10 plays a role in the modulation of neutrophil chemotaxis to FMLP and possibly IL-8. The observed shift in the FMLP dose response suggests that CD10 may exert its modulating effect on neutrophil responses to FMLP by cleavage of FMLP peptide bonds.

**Effects of interleukin 8 on neutrophil function and the interaction with FMLP**

**M MIKAMI, GG LLWELLYN-JONES, RA STOCKLEY Lung Immunobiochemical Research Laboratory, The General Hospital, Steelhouse Lane, Birmingham**

Interleukin 8 (IL-8) is an important inflammatory mediator which may play a role in the pathogenesis of lung diseases. However, there is some controversy regarding the effects of IL-8 on neutrophil function. It is known that IL-8 is a potent chemoattractant and is able to stimulate superoxide production, there is uncertainty whether it can alter neutrophil degranulation. Therefore we have designed experiments to look at the effect of IL-8 on resting neutrophils and their response to FMLP. IL-8 showed a dose response effect on chemotaxis with a maximum at a concentration of 5 nM (mean [SE] 22.7 (1.7) cells/field). Neutrophil degranulation was also increased from a control value of 1.93 (0.31) gN to 2.85 (0.33) gN at 10 nM (p<0.02). Finally, superoxide generation was increased from a control value of 2.79 (0.37) nmol/h to 4.31 (0.44) nmol/h (p<0.01) at 10 nM. Preincubation of neutrophils with IL-8 showed a dose dependent suppression of chemotaxis to FMLP from a control mean of 18.2 (4.2) to 8.5 (3.9) cells/field at 10 nM (p<0.01). When IL-8 was used as a chemoattractant together with FMLP it showed an additive effect at suboptimal concentration of each agent. However, at optimal concentrations the chemotaxis was increased less than would be expected for each alone (IL-8=17.1 (1-1) cells/field, FMLP=25.0 (2-6) cells/field, IL-8+FMLP=30.2 (2-6) cells/field). FMLP-induced neutrophil degranulation increased after preincubation of the cells with IL-8 (10 nM) from a mean of 4.67 (0.61) to 6.20 (0.77) gN (p<0.04). FMLP-induced superoxide generation also demonstrated an increase after preincubation with 5 nM IL-8 from a control mean of 15.58 (2.79) to 21.32 (2.1) nmol/h. This increase, 5.74 (0.94) nmol/h, was greater (p<0.01) than the effect of IL-8 treatment alone (increase = 0.84 (0.32) nmol/h), suggesting synergism. These data indicate that IL-8 is not only a potent chemoattractant for neutrophils but also that it stimulates other neutrophil functions such as degranulation and superoxide generation. Preincubation of neutrophils with IL-8 reduces the chemoattractant response to FMLP but has an additive effect with FMLP as a chemoattractant and stimulant of degranulation. Finally, IL-8 acts synergistically with FMLP for superoxide generation.

**Interleukin 8 may be involved in the recruitment of neutrophils into the respiratory tract of smokers and patients with chronic obstructive pulmonary disease**

**VM KEATINGS, PD COLLINS, DM SCOTT, TJ WILLIAMS, PJ BARNES Department of Thoracic Medicine and Applied Pharmacology, The National Heart and Lung Institute, London**

It is known that patients with chronic obstructive pulmonary disease (COPD) have increased numbers of neutrophils in the airway lining fluid compared with controls. This observation has also been made in smokers with normal lung function. The mechanisms for this are unknown. The cytokine interleukin 8 (IL-8) is produced by alveolar macrophages, epithelial cells, and neutrophils. Its production is stimulated by TNFα, interleukin 1, and neutrophil elastase. We wished to determine whether this cytokine is involved in the airway lining fluid neutrophilia of COPD. We induced sputum using inhaled hypertonic saline in 13 patients with COPD, 12 healthy smokers, and 16 normal non-smoking controls. Total and differential cell counts were carried out. A proportion of the sputum was ultracentrifuged at 60 000 g to obtain the cell phase which was assayed for IL-8 using a radioimmunoassay. Patients with COPD and healthy smokers had significantly higher IL-8 concentrations than the non-smoking controls (1.73, 0.57 and 0.13 nmol respectively, p=0.03 and p=0.02.) In the COPD group the IL-8 concentrations correlated strongly with the total neutrophil concentration (r=0.76), indicating that IL-8 and the neutrophilia are linked in this condition.

**Tumour necrosis factor α is present in high concentrations in the airways of patients with chronic obstructive pulmonary disease**

**VM KEATINGS, DM SCOTT, PJ BARNES Department of Thoracic Medicine, The National Heart and Lung Institute, London**

It is known that patients with chronic obstructive pulmonary disease (COPD) have increased numbers of neutrophils in the airway lining fluid compared with controls. This observation has also been made in smokers with normal lung function. The mechanisms for this are unknown. The cytokine tumour necrosis factor α (TNFα) is produced by alveolar macrophages and is chemoattractive for neutrophils both directly and through the induction of adhesion molecules. It has been implicated in the pathogenesis of the cellular infiltrate and the bronchial hyper-responsiveness seen in asthma. We wished to determine whether this cytokine was present in the airways of patients with COPD. We induced sputum using inhaled hypertonic saline in 13 patients with COPD, 13 patients with asthma, 12 healthy smokers, and 16 normal non-smoking controls. Total and differential cell counts were carried out. A proportion of the sputum was ultracentrifuged at 60 000 g to obtain the cell phase which was assayed for TNFα using an enzyme linked immunosorbent assay. Levels of TNFα did not differ between healthy smokers and normal controls (concentrations 26.5 and 16.3 pg/mL, respectively). Patients with COPD had significantly higher TNFα concentrations (98.1 pg/mL) than the smoking and non-smoking controls (p=0.04 and p=0.01). TNFα concentrations in the patients with COPD were higher than in the asthmatic group, but this did not reach significance (p=0.07). These data suggest that there is a significant inflammatory response in the airways of patients with COPD and that TNFα may play a significant part in this.

**Contribution of neutrophils, tumour necrosis factor and glutathione to cigarette smoke-induced increased airspace permeability**

**XY LI, IR HAHMAN, RK DONALDSON, WM MACNIE Respiratory Medicine Unit, Department of Medicine, University of Edinburgh; Department of Biological Sciences, Napier University, Edinburgh**

Increased airspace epithelial permeability (EP) is characteristic of cigarette smokers and is important in augmenting the inflammatory response in the airspaces and hence may have a role in the pathogenesis of emphysema. The purpose of this study was to investigate the mechanisms of this phenomenon. We have previously shown that intra-tracheal instillation of cigarette smoke condensate (CSC) induces increased EP in vivo in rats and in vitro in epithelial cell monolayers associated with a disturbance in glutathione (GSH), a major lung antioxidant. In this model neutrophils account for 5–10% of the bronchoalveolar (BAL) cells. Depletion of neutrophils and macrophages by an intratracheal injection of anti-neutrophil antibody did not influence the EP induced by CSC. The role of tumour necrosis factor (TNF) was also investigated. Although instillation of TNFa increased EP in the rat lung (0.62 (0.04) to 1.27 (0.29)%, p<0.001) 16 hours after instillation, only a trivial amount of TNF was detected in BAL fluid in vivo or in culture medium from BAL leucocytes obtained from CSC-treated animals. Furthermore, anti-TNF antibody did not abolish the increased EP induced by CSC. The CSC-induced increased EP was, however, associated with a profound fall in BAL GSH (2.56 (0.91))
to 0·31 (0·21–0·7 nmol/ml, p<0·001) and in lung GSH (869 (78) to 502 (70) nmol/g lung w.w., p<0·001) associated with an increase in oxidised GSH (90 (6) to 149 (85), p<0·01) one hour after instillation with a return to control values in lung but not in BAL six hours after instillation. These studies confirm an important role for the antioxidant glutathione in the increased epithelial permeability caused by cigarette smoke condensate.

Cigarette smoke and rat lung glutathione metabolism

I RAHMAN, X Y LI, K DONALDSON, W MAGNIE Respiratory Medicine Unit, Department of Medicine, University of Edinburgh; Department of Biological Sciences, Napier University, Edinburgh Cigarette smoke contains over 4700 highly electrophilic chemicals. The glutathione redox system is protective against smoke-induced injury. We studied the acute effects of cigarette smoke condensate (CSC) on glutathione (GSH) metabolism in rat lung and primary cultured type II alveolar epithelial cells. One hour after intratracheal instillation of CSC (table) and in vitro exposure of epithelial cells GSH diminished, without elevation of oxidised glutathione (GSSG) or protein-GSH mixed disulphides (PrSSG). However, the depletion of GSH was associated with rapid elevation of GSH conjugates unrelated to the action of glutathione-S-transferase. Activities of γ-glutamylcysteine synthetase, glutathione peroxidase, and glucose-6-phosphate dehydrogenase were significantly decreased after one hour, returning to normal levels after six hours exposure to CSC without change in glutathione-S-transferase and glutathione reductase activities. Thus CSC exposure causes transient depletion of GSH by the formation of GSH conjugates, in association with inhibition of GSH synthesis and other redox enzymes. These results are relevant to the changes in glutathione metabolism in smoker’s lungs and the associated airways injury. [Supported by the Tobacco Products Research Trust and The Norman Salvesen Emphysema Research Trust.]

<table>
<thead>
<tr>
<th>Time (hours)</th>
<th>GSH (nmol/g wet wt)</th>
<th>GSH conjugates (nmol/g wet wt)</th>
<th>GSSG (nmol/g wet wt)</th>
<th>PrSSG (nmol/g wet wt)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>780 (32)</td>
<td>22 (8)</td>
<td>94 (10)</td>
<td>32 (6)</td>
</tr>
<tr>
<td>1</td>
<td>510 (70)**</td>
<td>72 (7)**</td>
<td>105 (10)</td>
<td>28 (2)</td>
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<tr>
<td>6</td>
<td>770 (100)</td>
<td>44 (4)**</td>
<td>65 (4)</td>
<td>27 (2)</td>
</tr>
<tr>
<td>24</td>
<td>770 (45)</td>
<td>32 (8)</td>
<td>93 (5)</td>
<td>29 (2)</td>
</tr>
</tbody>
</table>

*p<0·05, **p<0·001. Values are mean (SE).