Written asthma action plans

M R Partridge

More widespread use of written asthma action plans should be encouraged

The first British guidelines for the management of asthma in adults published in 1990 clearly recommended self-management of asthma. The exact statement read: “As far as possible, patients should be trained to manage their own treatment rather than be required to consult their doctor before making changes”. Similar advice has been repeated in subsequent revisions of the UK guidelines and in the NHLBI global strategy for asthma management and prevention.

The evidence base for these recommendations is strong, and 36 trials comparing self-management education with usual care were reviewed for the Cochrane Library. This review suggested that self-management education could be associated with a reduction in hospital admissions of up to 40%, a reduction in emergency room visits of 20%, and similarly impressive reductions in unscheduled visits to the doctor, night time symptoms, and days off work or school. The authors concluded that training programmes that enabled people to adjust their medication using a written asthma action plan appeared to be more effective than other forms of self-management. In this issue of Thorax Gibson and Powell report the results of a further review to determine what is important about personalised written asthma action plans. They conclude that such plans are best when using 2–4 action points which involve increasing the dose of inhaled steroid and initiation of oral steroid therapy for exacerbations. Plans using peak flow should be specific barriers to implementation of such educational advice because of lack of awareness of the recommendations; erroneous belief that all asthma attacks are acute; lack of confidence in patients self-managing their own condition; dislike of self-management because (in some healthcare systems) it leads to loss of income; lack of physician confidence in teaching patients self-management skills; perceived lack of time.

Failure to implement recommendations contained within guidelines is, of course, not confined to failure to offer patients with asthma written personalised action plans. However, there may be specific barriers to implementation of such educational advice because of lack of specific training and knowledge regarding what should be given in the way of self-management advice and personal asthma action plans. Non-availability of partially preprinted material on to which advice may be written may also lead to patients not receiving such plans. Numerous studies have shown that, at least among adults, most asthma exacerbations, while often severe, are not acute. One study showed that 56% of adults admitted to hospital with severe asthma had experienced night time waking for at least 5 nights before admission. In another national census of those attending UK accident and emergency departments with asthma, one fifth of adults had been kept awake by their asthma for more than 3 nights before attendance. A study in Canada found that one fifth of patients with asthma admitted to hospital and one fifth of those requiring intensive care had had symptoms for at least 21 days before admission. These studies suggest that, for most patients with troublesome asthma, plenty of time was available for either the patient or the doctor to alter treatment to avoid deterioration to the point where the patient needed to be admitted to hospital.

“There can be no further excuse for delaying widespread implementation of . . . written personal asthma action plans”

In some healthcare systems the concept of devolving care to the patient may have negative financial implications for health professionals. This might lead to them being reluctant to implement recommendations regarding the issuing of personal asthma action plans. It would be a pity if the beneficial results from 36 good clinical trials were to be negated by such financial considerations. Perhaps such colleagues could be convinced of the advantages of a partnership approach to medical care by pointing out that, in another study, over 30% of patients who scored their physicians as being “non-participatory” changed physicians over the subsequent year, whereas those who scored their physicians as being “participatory” were half as likely to report that they would change their physician in the following 12 months.

Time is needed to teach patients how to recognise signs of deteriorating asthma and to teach self-management skills, but Clark and colleagues have shown that such training, when offered in the context of an interactive educational seminar, can have a lasting effect on physician behaviour and better outcomes, and consultation times are not necessarily extended. In some healthcare systems such tasks are helpfully shared with nursing colleagues.

In their paper in this issue of Thorax, Gibson and Powell emphasise that action plans which involve both increasing inhaled steroid dosage and taking steroid tablets are the most effective, yet some may perceive a controversy with regard to increasing the dose of inhaled steroids. The British asthma guidelines...
state that the value of doubling the dose of inhaled steroids is unproven. However, this statement must be seen within the context of most of the published studies of self-management education which have included a description of the action plans used, recommending a doubling or trebling of the dosage of inhaled steroids. This paradox can be explained either by understanding that advice to double the inhaled steroids is only effective if given within the wider context of self-management education, or by an appreciation that it is the concept of varying dosage of medications that is important rather than the actual magnitude of change. It may be that the advice in zone 2 of a personalised asthma action plan also works by reminding the non-compliant patient to take his or her inhaled steroid. A further explanation is that doubling alone may not be sufficient. An Italian study suggested that the most efficacious interventions were probably those which involved reducing the dose of inhaled steroid when well controlled and then quadrupling it at the first sign of loss of control of asthma.16 The need for us to teach patients how to both increase and decrease their dose of inhaled steroids is exemplified by recent work which showed that, in a group of adults with asthma, the active practice of stepping down treatment was associated with a mean reduction in daily inhaled steroid usage of 348 μg beclometasone equivalent per day, with equally good outcomes to those who had stayed on a fixed dose.17

Patients with many long term conditions fail to comply with their therapeutic regimen. Many understandably express negative views regarding their “dependence” on regular medication, and many patients stop, start, or vary the dose of their medication. Self-management education and the issuing of personalised written action plans permit us to hand control of their condition to patients in such a way that they vary their dose of medication in a scientific manner, rather than according to whim. Patients dislike the uncertainty associated with a variable condition such as asthma, and they dislike dependency upon health professionals.

Teaching patients how to vary their dose of asthma medication according to their symptoms and according to the severity of their condition returns control to them and has been shown to be associated with enhanced compliance.18 Health service usage by those with asthma is reduced by such actions and it is likely that, overall, there may be a reduced usage of medication and financial benefits.19 There can be no further excuse for delaying widespread implementation of the issuing of written personal asthma action plans. Gibson and Powell have given us a clear steer as to the important constituents of such plans. Further research is still needed into which subgroups of those with asthma benefit most, and why, for some such as the repeatedly hospitalised,20 such benefit is less.

How we encourage self-management education and the issuing of personalised asthma action plans by health professionals is similarly unclear.21 Part of the failure may have been our failure to teach better what is involved or may reflect poor availability of materials. Part may reflect lack of motivation of healthcare professionals or lack of time. It will be of interest to see how much the financial incentive of the asthma 3+ visit plan in primary care improves implementation in Australia.22


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COPD in the ECRHS

J Vestbo

More work is needed on the concept of staging of COPD

As a junior doctor I once worked in a hospital where the leading consultant in medicine refused to accept the diagnosis of asthma in patients older than 40 years. To him, airflow obstruction was “asthma” in the young and “chronic bronchitis” in the elderly. While it soon became apparent that asthma does occur after the age of 40, the likelihood of significant airflow limitation occurring in young adults who have never had asthma has always seemed small to me. In this issue of Thorax De Marco et al describe the prevalence of chronic obstructive pulmonary disease (COPD) in young adults taking part in the European Community Respiratory Health Survey (ECRHS). They found COPD to be a considerable issue; in total, 3.6% had COPD stage I+, according to the NHLBI/WHO Global Initiative for Chronic Obstructive Lung Disease (GOLD).11 12 and 11.8% had chronic respiratory symptoms without airflow limitation—that is, COPD GOLD stage 0.

The study raises several questions relating to methodology, findings, and interpretation. Diagnosis and staging of COPD was done according to the GOLD guidelines12 using an FEV1/FVC ratio of 0.7 and FEV1 cut off points of 80%, 50%, and 30%. In subjects aged 20–44 years a ratio of 0.7 will not overestimate airflow obstruction—more likely it will underestimate it. The major challenge seems to be exclusion of asthma and the approach of De Marco et al can, to some extent, be questioned. In contrast to GOLD recommendations, prebronchodilator FEV1 was used for staging but this seems acceptable in the epidemiological setting where administration of bronchodilators is often not feasible. Patients with self-reported asthma without cough/phlegm were excluded while those with both self-reported asthma and chronic symptoms were considered to have COPD with coexisting asthma. The latter seems intuitively correct in a 44 year old heavy smoker with a smoking history of 30 pack years, but is it true in the 20 year old never smoker with self-reported asthma? Unfortunately, no valid answers exist; GOLD has not attempted to separate stage 0 COPD from symptomatic asthma, and only for subjects with irreversible airflow limitation does GOLD acknowledge the problem: “Poorly reversible airflow limitation associated with bronchiectasis, cystic fibrosis, tuberculosis, or asthma is not included except insofar as these conditions overlap with COPD.”7 In the Copenhagen City Heart Study cohort13 54% of women and 63% of men with self-reported asthma had chronic productive cough; this will presumably remain an issue for debate for some time.

COPD stage 0 denoting subjects “at risk” was introduced by GOLD, but the concept cannot be regarded as evidence based and remains controversial.14 It is, nevertheless, intriguing that the prevalence of chronic symptoms in 20–44 year old subjects is more than 10% on average and as high as 24% in Spain. Risk factors did not differ substantially between stages 0 and 1+, and a recent Italian study has shown that stages 0 and 1 differ little in health status.8 Still, we do need prospective studies of stage 0 including various outcomes. We also have to make clear the reason for applying staging to COPD. Undoubtedly, staging facilitates communication and comparison of study results. It is, however, less clear that it reflects biological changes over time. The concept of cancer staging—where, by definition, patients progress through the stages—may not be valid in COPD. While it is unlikely for anyone to have stage III or IV without passing through earlier stages, COPD stage I can undoubtedly develop without the patient ever having been in stage 0.9 Years of looking at the “Fletcher diagram”10 have anchored the impression of rapid decline so firmly in our minds that we may tend to forget that, through impaired growth of lung function in childhood and early adolescence, any superimposed airflow obstruction at a later age could very well start the patient off in COPD stage II.11 For this and other reasons, more work on the concept of staging of COPD is clearly needed.

COPD is a burden in the elderly, but it is not a disease of the elderly alone. The notion of COPD in young adults was confirmed by the “confronting COPD” study, but whereas that study used doctor’s diagnosis and presence of symptoms, the ECRHS study has verified the diagnosis with spirometric testing in random population samples, enabling us to quantify the problem. Unfortunately, the study by De Marco et al does not tell us the prevalence of doctor diagnosed COPD in their cohort. COPD is often undiagnosed10 and, based on data from the IBERPOC study, this is even more so in younger patients11 and in women more than in men.11 12 In this respect, the ECRHS study showed COPD to be more prevalent in men than in women. When biological explanations are applied to these findings, caution is probably warranted. Better information is available in this area from longitudinal studies13 and, in addition, detailed information on smoking such as age of starting and inhalation is essential for adjusting properly for sex differences in smoking habits when addressing susceptibility.14

With the study by De Marco et al, however, COPD epidemiologists now have to join asthma epidemiologists in praising the ECRHS. One important question remains: How should these findings change our perception of COPD? They probably should not! The strengths of the paper lie in the finding that COPD is a widespread problem in young adults and the implications of the quantification. To limit case finding and/or screening for COPD to middle aged or elderly subjects would be missing a window of opportunity based on these findings.

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Oxygen sensing

New insights into oxygen sensing at a cellular level

S R Walmsley, K K K Sheares, A Sobolewski, N W Morrell, E R Chilvers

The oxygen sensing pathway offers a new set of therapeutic targets for conditions ranging from inflammatory lung disease to pulmonary hypertension

The ability of cells to detect and respond to a fall in oxygen tension is of fundamental importance for maintaining oxidative metabolism and tissue homeostasis. One of the challenges facing scientists working in this area has been that any proposed mechanism for oxygen sensing has to accommodate the very different tolerances of certain tissues to hypoxia and the extreme variation in the cellular responses observed. Hence, while skeletal muscle cells can recover function after 30 minutes of anoxia, the brain suffers irreparable damage after only 4–6 minutes of ischaemia. Moreover, while carotid body cells respond to changes in oxygen tension that barely register in non-chemosensory tissues (and do so within seconds), upregulation of erythropoietin synthesis in the interstitial peritubular cells is transcriptionally regulated and requires far more protracted periods of hypoxaemia. Despite such variances in oxygen sensitivity and response time, all cells appear capable of responding to hypoxia and the essential components of a universal oxygen sensing mechanism have at last begun to emerge. Moreover, from studies conducted in stroke and heart disease, it is apparent that therapeutic targeting of this novel pathway is set to transform our approach to pathology previously deemed intractable.

RESEARCH STUDIES

Initial clues into how cells respond to low oxygen came from studies undertaken in the early 1990s examining the hypoxic response element (HRE) of the erythropoietin (Epo) gene. This led to the identification of a transcriptional activator called hypoxia inducible factor (HIF). HIF is a heterodimer composed of HIF-1β (or aryl hydrocarbon receptor nuclear translocator, ARNT), which is constitutively expressed, and HIF-1α whose expression and transcriptional activity are tightly regulated by the ambient oxygen concentration. Once formed, this protein complex migrates to the cell nucleus and, together with the co-activator CBP/p300, binds to the HREs present on the promoter region of genes involved in regulating metabolic supply and demand. Examples of HIF regulated genes include those involved in regulating vascular tone (for example, iNOS and adrenomedullin), angiogenesis (for example, vascular endothelial growth factor, VEGF), cell metabolism (for example, lactate dehydrogenase A, the glucose transporter GLUT-1), and haemoglobin biosynthesis (for example, erythropoietin). These findings, together with the ubiquitous nature of HIF and the demonstration that HIF deficient animals show major defects in many core physiological responses to oxygen, have resulted in HIF being regarded as one of the master regulators of the cellular response to hypoxia.

The next step in this quest was to define the mechanism responsible for hypoxic induction of HIF-1α. Through a combined structural and genetic approach, it has now been possible to show that HIF-1α activity is regulated by enzymatic hydroxylation at specific prolyl and asparaginyl residues by a novel 2-oxoglutarate dependent class of oxygenases. Critically, these enzymes have been shown to display an absolute requirement for oxygen in addition to iron (Fe²⁺) and ascorbate. These oxygen sensitive enzymes inhibit HIF activity in a complementary manner since the prolyl hydroxylase domain containing enzymes (PHDs) result in an interaction between HIF-1α and the von Hippel-Lindau protein which targets HIF-1α for proteosomal destruction, and factor inhibiting HIF (FIH) causes asparaginyl hydroxylation and blocks HIF association with CBP/p300. Hence, under normoxic conditions, HIF-1α levels remain low and this prevents the transcription of genes containing HRE promoters (fig 1).

CLINICAL APPLICATIONS

How does this inform our understanding of the pathophysiology of lung disease and can it provide the basis of novel therapies? Mice homozygous for a null mutation in the HIF-1α or HIF-1β genes die at mid gestation with vascular defects primarily involving the embryonic and extraembryonic circulation, respectively. In contrast, HIF-1α−/− mice develop normally and are indistinguishable from wild type littermates. However, when exposed to 10% oxygen for up to 6 weeks, the HIF-1α−/− mice demonstrate reduced susceptibility to pulmonary hypertension, polycythaemia, and right ventricular hypertrophy relative to their wild type littermates. In the lungs of HIF-1α−/− mice, the expression of genes containing HREs is lower than in wild type littermates. Morphometric analysis showed that the chronically hypoxic HIF-1α−/− mice have fewer completely muscularised pulmonary arterioles and the degree of muscularisation in such vessels is reduced compared with HIF-1α wild type mice. Thus, HIF-1 appears to play a major role in mediating pulmonary vascular remodelling in chronic hypoxia, and therapeutic manoeuvres that inhibit HIF-1 activity in the lung may slow the progression of chronic pulmonary hypertension.

Abbreviations: ARNT, aryl hydrocarbon receptor nuclear translocator; FIH, factor inhibiting HIF; HIF, hypoxia inducible factor; HRE, hypoxic response element; PHD, prolyl hydroxylase domain containing enzyme.

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7 Prescott E, Oster M, Veselka J. Importance of initial clues into how cells respond to a fall in oxygen tension to show that HIF-1α null mutation in the HIF-1α embryonic and extraembryonic circula- tion, respectively. In contrast, HIF-1α−/− mice develop normally and are indis- tinguishable from wild type litters. However, when exposed to 10% oxygen for up to 6 weeks, the HIF-1α−/− mice demonstrate reduced suscepti- bility to pulmonary hypertension, polycythaemia, and right ventricular hyper- trophy relative to their wild type litters. Morphometric analysis showed that the chronically hypoxic HIF-1α−/− mice have fewer completely muscularised pulmonary arterioles and the degree of muscularisation in such vessels is reduced compared with HIF-1α wild type mice. Thus, HIF-1 appears to play a major role in mediating pulmonary vascular remodelling in chronic hypoxia, and therapeutic manoeuvres that inhibit HIF-1 activity in the lung may slow the progression of chronic pulmonary hypertension.

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hypoxia and the firing of carotid body glomus cells. The current contenders in this area are the mitochondrial NAD(P)H oxidases, which are postulated to produce a diffusible redox mediator in response to hypoxia.

CONCLUSIONS

These studies show that, while cells display widely differing tolerances and responses to hypoxia, oxygen sensing by dioxygenases and their consequent effects on HIF dependent transcriptional events appear to be a property that is shared by most, if not all, cells. This pathway now offers a new set of therapeutic targets for an array of lung diseases ranging from inflammatory lung disease through to pulmonary hypertension.

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22 Ehrenreich H, Hasselblatt M, Dembowski C, et al. Erythropoietin therapy as a number of interventional available are beneficial to produce a beneficial outcome.5–11 For example, population screening studies suggest that individuals identified as having PI*ZZ AAT deficiency at birth are less likely to start smoking or to continue smoking than age matched adolescent peers.5 10 Furthermore, although no definitive randomised clinical trial has established the clinical efficacy of augmentation therapy—currently the intravenous infusion of purified pooled human plasma $\alpha_1$-antitrypsin—the results of many observational studies strongly suggest its clinical efficacy.9 10 15 Indeed, a recent standards document from the Canadian Thoracic Society and the aforementioned evidence-based standards document which is supported and endorsed by the European Respiratory Society, the American Thoracic Society, the American College of Chest Physicians, the American Respiratory Care Foundation, and the Alpha-1 Foundation both conclude that augmentation therapy has clinical efficacy.17 As reviewed in this series of articles, new and emerging therapies offer the prospect of even more effective treatments that can be made available to individuals known to have severe AAT deficiency.18
It is hoped that this comprehensive review will enhance the recognition of

$\alpha_1$-Antitrypsin deficiency

A new series focusing on this important and under-recognised illness

$\alpha_1$-Antitrypsin deficiency

J K Stoller

A $\alpha_1$-antitrypsin (AAT) deficiency is a common but under-recognised clinical entity.1–3 The editors of Thorax have therefore commissioned a series of papers by internationally recognised experts on the key clinical and investigative concepts in this important disease, which will offer the reader an up to date summary of AAT deficiency. Topics to be addressed include:

- the epidemiology of AAT deficiency;
- genetic aspects of AAT deficiency: phenotypes and genetic modifiers of emphysema;
- clinical manifestations and natural history of AAT deficiency;
- molecular pathophysiology of AAT deficiency;
- pathogenesis of lung disease in AAT deficiency;
- intravenous augmentation therapy for AAT deficiency: current understanding;
- new and emerging therapies for AAT deficiency; and
- CT imaging in AAT deficiency.

Why this attention to AAT deficiency now? As mentioned above, despite the fact that it affects up to one in 1600 newborn infants,2 AAT deficiency is both under-recognised and “under-understood”.4 As evidence of this under-recognition, in a survey of 300 PI*ZZ individuals the mean interval between the appearance of the first attributable symptom (almost invariably dyspnoea due to fixed airflow obstruction) and the diagnosis of AAT deficiency in a group of mean age 49 years was 7.2 years.5 Furthermore, 44% of the respondents reported seeing at least three physicians before the diagnosis of AAT deficiency was made. In the United States, of the expected 80 000–100 000 with severe AAT deficiency (PI*ZZ), fewer than 10% have been clinically recognised.6

Since establishing the diagnosis is both easy (a simple blood test for a serum level and, if low, a phenotype to secure the diagnosis) and relatively inexpensive (less than $US200 for a serum level or a phenotype in most commercial laboratories and widespread availability of free testing services), there are ample opportunities to increase recognition of the condition by enhancing clinicians’ diagnostic suspicion of AAT deficiency.

What clinical features should clinicians look for? As discussed in this series and in a recently published evidence-based standards document on the diagnosis and management of individuals with AAT deficiency,7 AAT deficiency should be suspected in patients with fixed airflow obstruction (especially in the absence of cigarette smoking or predisposing occupational exposures), in individuals whose family history suggests emphysema and/or liver disease, and in those with suggestive clinical characteristics—for example, basilar hyperlucency on the chest radiograph, bronchiectasis that is otherwise unexplained, panniculitis, or cirrhosis for which a known aetiology such as viral hepatitis, haemochromatosis, or Wilson’s disease is not evident.

Why make the diagnosis of AAT deficiency? As with all diseases, the impetus to diagnose is the desire to ameliorate the adverse effects of the illness, to prolong life, to improve the quality of life of affected individuals and, in the case of a genetic disease like AAT deficiency, to counsel at risk family members in order to encourage health attentive behaviour and treatment that will lessen the likelihood of disease and/or forestall its progression. Indeed, it is important to establish the diagnosis of AAT deficiency as many interventions are available that can produce a beneficial outcome.5–11 For example, population screening studies suggest that individuals identified as having PI*ZZ AAT deficiency at birth are less likely to start smoking or to continue smoking than age matched adolescent peers.5 10 Furthermore, although no definitive randomised clinical trial has established the clinical efficacy of augmentation therapy—currently the intravenous infusion of purified pooled human plasma $\alpha_1$-antitrypsin—the results of many observational studies strongly suggest its clinical efficacy.9 10 15 Indeed, a recent standards document from the Canadian Thoracic Society and the aforementioned evidence-based standards document which is supported and endorsed by the European Respiratory Society, the American Thoracic Society, the American College of Chest Physicians, the American Respiratory Care Foundation, and the Alpha-1 Foundation both conclude that augmentation therapy has clinical efficacy.17 As reviewed in this series of articles, new and emerging therapies offer the prospect of even more effective treatments that can be made available to individuals known to have severe AAT deficiency.18

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individuals with AAT deficiency by clinicians and foster optimal medical management.


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