Implementation of British Thoracic Society guidelines for acute exacerbation of chronic obstructive pulmonary disease: impact on quality of life

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BACKGROUND: The British Thoracic Society (BTS) guidelines have not been examined collectively for their impact on chronic obstructive pulmonary disease (COPD). Whether intensive outpatient follow up of COPD patients after acute admission, using these guidelines, improved quality of life compared to the "usual practice" of primary care follow up was investigated.

METHODS: Altogether 103 patients with a new diagnosis of COPD were admitted and screened over a four year period. Seventy patients were excluded because of another dominant medical condition or a mandatory requirement for intervention. Patients were randomised to regular primary care (control group, n = 15) or chest clinic follow up (intervention group, n = 10). Spirometry, oxygen saturation, St George’s Respiratory Questionnaire (SGRQ), and Short Form 36 questionnaire were measured at baseline and six months. The intervention group was reviewed at least four times in the six month period and received spirometry, ambulatory oxygen assessment, smoking cessation advice, nebuliser assessment, a steroid trial, advice about nutrition/exercise, and introduction to a patient support group.

RESULTS: There was no significant difference between baseline measurements in the two groups. There was a significant mean (SD) improvement in the SGRQ symptom score from baseline to six months in the intervention group [20.98 (20.36)] compared with the controls [0.23 (12.55)] (p = 0.004). At six months the SGRQ symptom score, impact score, and total score was significantly better in the intervention than the control group (p=0.01, 0.02, and 0.02).

CONCLUSION: Aggressive implementation of BTS guidelines after initial hospitalisation may improve respiratory health specific quality of life scores in patients with COPD. Larger studies are needed to confirm this finding.
Ethical approval
The Redbridge and Waltham Forest Research Committee gave ethical approval for this study.

RESULTS
Baseline data
Of the 103 patients screened, 70 did not satisfy the entry criteria, predominantly because of need for mandatory intervention or severe co-morbidity, and were excluded from the study. The remaining 33 patients were randomised to regular chest clinic follow up (intervention group) or primary care follow up (control group). Eight patients (five from the intervention and three from the control group) withdrew from the study resulting in 15 patients in the control group and 10 in the intervention group who completed the study. Patients withdrew from the control group because one patient was lost to follow up and two had repeated admissions and from the intervention group because two patients were lost to follow up, one inadvertently attended their general practitioner for follow up, and two patients had repeated hospital admissions. There were no significant differences between the two groups (control v intervention) in the baseline quality of life scores, age (71.33 (8.39) v 68.40 (5.76) years), FEV₁ (39 (11.88) v 44.70 (21.84) percent predicted), and severity of disease according to the BTS guidelines for COPD (20% mild, 27% moderate, and 53% severe disease v 20% mild, 20% moderate, and 60% severe disease).

Symptom scores at six months
The mean improvement of the SGRQ symptom score from baseline to six months in the intervention group (20.98 (20.36)) was significantly better compared with the control group (0.23 (12.55); p = 0.004) (fig 1). In the intervention group, at six months the SGRQ symptom score, impact score (psychosocial function), and total score was significantly better than the control group (p = 0.01, 0.02, and 0.02) (fig 2). Although there was a trend towards improvement in the Short Form 36 general health score (p = 0.067), this did not reach statistical significance.

There were fewer readmissions (2 v 9) and patients with two or more exacerbations within a six month period (2 v 3) in the intervention group but the small numbers precluded meaningful statistical analysis. There was no significant difference at six months in FEV₁, resting oxygen saturations and the amount of oxygen, nebuliser, theophylline, and long acting bronchodilator prescriptions in the two groups.

DISCUSSION
This study suggests that regular follow up in the outpatient department accompanied by aggressive implementation of BTS management guidelines produces disease specific improvements in quality of life indicators but not in general health quality of life. Despite the enormous disease burden of COPD within the UK there are few studies addressing the value of the BTS recommendations for active follow up of admitted patients. Indeed most of the intervention suggestions are individually unproven in this area and certainly have not been studied as a combined package of measures. The notable exception is the literature on pulmonary rehabilitation and this study purposefully excluded that intervention of proven value.

This current study is weakened by the small number of patients who completed the study. An important lesson can be learned from this. The small number enrolled was mainly due to the disease severity of the admitted patients that meant they could not ethically be randomised to the control group—for example, investigation of possible lung cancer and need for long term oxygen therapy. Others had severe co-morbidity that dominated their illness profile and meant that COPD interventions may be ineffective. Finally, of those recruited a significant number withdrew because of concurrent illness and nine of 15 patients in the control group were readmitted within the six months of follow up period concordant with other research in this area.5

The Short Form 36 scores did not improve to the same extent that the SGRQ scores did. This discrepancy may be explained because the Short Form 36 is a more generic measure while the SGRQ is a more disease specific measure, with a much higher proportion of its content directly relevant to COPD.7

The observed change in the SGRQ could not be accounted for by changes in drug prescriptions or lung function. This was not surprising as the degree of correlation between objective measures of lung function and quality of life measures is inconsistent.8–10 As the symptoms of COPD are complex and multicausal it is possible that the secondary care package may have made an impact on aspects of COPD other than lung function. It may be that the simple effect of regular contact with a healthcare professional is beneficial and it remains unproven if any of the interventions made altered the pathophysiology of the COPD process or if simply the whole package of care provided a placebo impact.

This report is noteworthy for two reasons. Firstly it objectively justifies the consensus derived recommendations for a package of measures to be allocated to patients discharged after a first admission with COPD. Such an allocation of secondary care resources would, however, be subject to a consideration of the cost effectiveness of this approach, an aspect not examined in this study. Secondly it highlights the surprising paucity of literature on this important subject. Larger multicentre studies are needed to confirm our findings and to identify the individual interventions that may account for a change in quality of life indices and the mechanisms by which this is achieved.
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Conflicts of interest: Dr G Hagan worked as an Honorary Clinical Assistant at Whipps Cross Hospital while in the employ of Glaxo Wellcome at the time of the study.

REFERENCES


IF ONLY I’D KNOWN

Ten things I wish I’d known when I was 25

(1) That everyone who tries to live forever eventually dies in the attempt.
(2) That older folk—particularly those in charge—don’t do what they do simply because they have lost their marbles along the way.
(3) That giving patients what they want rather than what they really need will save you a lot of heartache.
(4) Save your energy only for the few battles you could actually win.
(5) Find out as soon after the start of the consultation begins as possible what the patient wants from you—it’s rarely what you imagine they should want.
(6) Very rarely is the patient’s view of themselves or their problem anywhere near shared by what those who know them well. Canvass a variety of views before accepting the patient’s version.
(7) That excellence in academic performance is rarely rewarded with popularity—but is worth it anyway.
(8) That in a profession where salaries are all the same—relatively small differences in advantage seem to matter hugely and cause massive envy. Try to hide your success as much as possible, or if uncovered, deny it.
(9) That there is much more to death than medical failure and that there is much more to life than being a doctor.
(10) That doctors not only alarmingly rarely really know what’s best for the patient—they don’t know what’s best for doctors—take all career advice with a pinch of lithium.

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