Effect of ethnicity on the prevalence, severity, and management of COPD in general practice

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INTRODUCTION

In the UK, chronic obstructive pulmonary disease (COPD) accounts for 30,000 deaths a year, 1.4% of GP consultations, 2% of hospital admissions, and costs the NHS £800 million. The main cause of COPD is tobacco smoking, but it is also associated with other factors such as occupational exposure, urban living, alpha-1 antitrypsin deficiency, and childhood respiratory infections. The majority of routine COPD management takes place in primary care, with the most effective intervention being smoking cessation. The COPD guidelines from the National Institute for Health and Clinical Excellence (NICE) recommend a stepwise approach to COPD prescribing and a range of non-pharmacological interventions. The Quality and Outcomes Framework (QOF) data from general practice estimates the UK COPD prevalence to be 1.4%, but epidemiological studies suggest the prevalence is considerably higher—a cross-sectional study using data from the Health Survey for England estimated the UK COPD prevalence to be 3.1% in 2005. COPD is more prevalent in males, older people, and populations that are deprived and urban. There is little data on the variation in UK COPD prevalence by ethnicity; Nacul et al found high COPD prevalence in black males and low COPD prevalence in South Asian females in the UK, but did not adjust for deprivation. In the US, Chatila et al found that African Americans presented with COPD at a younger age than white populations, despite smoking less. However, in a different US study, Vollmer et al found no significant difference in COPD prevalence by ethnicity. COPD prevalence is known to be strongly associated with smoking prevalence. The 2004 Health Survey for England found that males from minority ethnic groups were more likely, and females less likely, to smoke than the general population.

This study was set in three contiguous inner east London primary care trusts (PCTs) where 53% of the population are members of minority ethnic groups; this is a much higher proportion than in the whole UK population, in whom minority ethnic groups total 7.9%. The populations of these three east London PCTs are among the eight most deprived localities in Britain. The aim of this study was to examine the effect of ethnicity on the prevalence, severity, and management of COPD.

METHOD

A total of 140 general practices in east London contributed routinely collected data as part of a regular annual audit of chronic disease management. These practice data cover 98% of the GP-registered population in the three PCTs [920,000 in February 2011].

Clinical data collection

Practice computer databases were interrogated using Egton Medical Information Systems web software. As Read Codes are the clinical classification system

Abstract

Background

Chronic obstructive pulmonary disease (COPD) remains a major cause of mortality and hospital use. Little is known in the UK about the variation in COPD prevalence, severity, and management depending on ethnicity.

Aim

To examine differences by ethnicity in COPD prevalence, severity, and management.

Design & setting

Cross-sectional study using routinely collected computerised data from general practices in three east-London primary care trusts (Newham, Tower Hamlets, and City and Hackney) with multiethnic populations of people who are socially deprived.

Method

Routine demographic, clinical, and hospital admission data from 194 practices were collected.

Results

Crude COPD prevalence was 0.9%; the highest recorded rates were in the white population. Severity of COPD, measured by percentage-predicted forced expiratory volume in 1 second, did not vary by ethnicity. South Asians and black patients were less likely than white patients to have breathlessness, indicated by a Medical Research Council dyspnoea grade of ≥4 (odds ratio [OR] 0.7 [95% confidence interval (CI) = 0.6 to 0.9]) and 0.6 (95% CI = 0.4 to 0.8)). Black patients were less likely than white patients to receive inhaled medications. Influenza and pneumococcal vaccine rates were highest among groups of South Asians (OR 3.0 [95% CI = 2.1 to 4.3]) and 1.8 [95% CI = 1.4 to 2.3]) respectively. Both minority ethnic groups had low referral rates to pulmonary rehabilitation. In Tower Hamlets, black patients were more likely to be admitted to hospital for respiratory causes.

Conclusion

Differences in COPD prevalence and severity by ethnicity were identified, and significant differences in drug and non-drug management and hospital admissions observed. Systematic ethnicity recording in general practice is needed to be able to explore such differences and monitor inequalities in healthcare by ethnicity.

Keywords

chronic disease management; chronic obstructive pulmonary disease; ethnicity; general practice; health inequalities.
used in UK general practice, all adult patients aged ≥35 years with a computerised diagnostic Read Code for COPD were included.15 Data was anonymised and managed according to NHS information governance requirements.16 Variables included in the analysis were recorded between 1 January 2009 and 31 December 2010. Clinical data included percentage-predicted forced expiratory volume in 1 second (FEV1%) — a low FEV1% denotes poor lung function and more severe COPD. The latest NICE guidance groups FEV1% into the following categories: very severe (<30); severe (30–49); moderate (50–79); and mild (≥80). The Medical Research Council (MRC) dyspnea scale indicates the level of symptoms related to breathlessness reported by the patient; higher scores indicate worse symptoms of breathlessness. Other clinical variables included: smoking status (grouped into current smoker and non-current smoker, including ex-smokers); blood pressure; total cholesterol; body mass index (BMI); and diagnosis of ischaemic heart disease, hypertension, diabetes, and asthma. Process of care variables included: pneumococcal immunisation, influenza immunisation, referral to pulmonary rehabilitation, and an annual COPD review. Prescription data included short-acting beta-agonists (SABAs), short-acting muscarinic-antagonists (SAMAs), long-acting beta-agonists (LABAs), long-acting muscarinic-antagonists (LAMAs), inhaled corticosteroids (ICS), combined inhaled corticosteroids/long-acting beta-agonists (ICS/LABAs), and theophylline. Hospital admissions (Tower Hamlets PCT population only) were collected using healthcare resource group categories. For each patient, a measure of all hospital inpatient episodes and respiratory specific episodes was included.

Ethnicity and socioeconomic variables
Self-reported ethnicity was recorded at the practice during registration or routine consultations. Ethnic groups were based on the UK 2001 Census and, for this study, were collapsed into five categories:
- white (British, Irish, other white);
- black (black African, black Caribbean, black British, other black, and mixed black);
- South Asian (Bangladeshi, Pakistani, Indian, Sri Lankan, British Asian, other South Asian, or mixed Asian);
- Other (Chinese, other ethnic groups, other mixed groups); and
- Unknown.

How this fits in
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Statistical analysis
All statistical analyses used Stata (version 9, StataCorp). In descriptive statistics, categorical variables were assessed for differences by using a χ2 test; continuous numerical variables were assessed using analysis of variance.

Differences in the prevalence and severity of COPD between ethnic groups were expressed as odds ratios (ORs) using stepwise logistic regression, with white ethnicity considered as neutral risk (OR 1.0). Analyses were adjusted by patient characteristics (age, sex, Townsend score, PCT, and asthma diagnosis, and clustered by general practice). The MRC dyspnoea scale was also adjusted by the comorbidity of hypertension, ischemic heart disease, and diabetes. Drug prescribing, non-pharmacological managements, and hospital admission rates were also adjusted by the FEV1% severity indicator.

RESULTS
Of the 7901 cases of GP-recorded COPD, ethnicity was recorded for 98.7% and
spirometry values in the previous 2 years were available for 5471. There was no significant difference in missing spirometry values between the three ethnic groups.

Crude COPD prevalence in the east-London population is 0.9%. (1.5% when adjusted to the European standard population. The recorded COPD prevalence for those aged ≥35 years in the study population was 2.5%. The highest prevalence was in the white population (4.4%); prevalence in the South Asian group was 1.3%, and the lowest rates were recorded in the black population (0.6%) (Figure 1). Figure 1 also shows that the white population had the highest prevalence of COPD in all age groups.

The prevalence of COPD by sex varied greatly by ethnicity. In the white population, 48.2% were male; in contrast, 81.2% and 64.4% were male in the South Asian and black populations respectively.

The major risk factor for COPD is smoking. The prevalence of GP-recorded current smoking rates in the adult population for the main ethnic groups in this study is shown in Figure 2. Smoking prevalence was 13.1% among those aged ≥25 years. The white population had the highest smoking prevalence at 18.8%; the prevalence among South Asians was 10.8% and 11.2% in the black population.

Smoking patterns by sex varied between ethnic groups and reflected the sex distribution of COPD prevalence. Among white smokers, 64.2% were male, among the South Asian group 88.7% were male, and in the black population 73.0% were male (data not shown).

Smoking prevalence by ethnicity was then examined within the COPD population. Table 1 illustrates that South Asian and black patients with COPD are much less likely to be current smokers than white patients. This suggests that these groups are more likely to stop smoking when there is a diagnosis of COPD.

The severity of COPD by ethnicity was examined using both FEV1% values and MRC breathlessness scores (Table 2). Using logistic regression, no difference was found in FEV1% by ethnicity when adjusting for demographic variables, Townsend score, PCT, and asthma, and clustering by practice. However, compared with the white population, both South Asian and black populations were less likely to have an MRC grade of ≥4, OR 0.6 (95% confidence interval [CI] = 0.4 to 0.8) and 0.7 (95% CI = 0.6 to 0.9) respectively; this suggests less severe symptomatic breathlessness in these groups, compared with white patients.

Drug prescribing and non-pharmaceutical management

Table 3 shows, by ethnicity, the prevalence of prescribing for each major COPD drug treatment type and key non-pharmaceutical management interventions. At the time of the study, only 56.3% of the whole COPD population was on a SABA; in contrast, more than three-quarters of the cohort was on either ICs alone (27.3%) or ICs combined with a LABA (51.9%). Less than a fifth of patients with COPD had a record of referral to
The adjusted analysis in Table 4 shows that black patients are less likely to receive SABAs, SAMAs, LAMAs, combined ICs/LABA, and theophylline, compared with their white counterparts. South Asians are less likely to receive LAMAs (adjusted OR 0.8 [95% CI = 0.6 to 0.9]) than their white counterparts. No difference was found for LABAs and ICs prescribed separately, by ethnicity.

Table 4 shows that among the non-pharmacological interventions, South Asians are more likely to receive both influenza and pneumococcal immunisations compared with white or black populations. Both South Asian and black patients are less likely to receive pulmonary rehabilitation referral compared with white patients.

**Hospital admissions**

Due to technical limitations, hospital admissions data were only available for the Tower Hamlets PCT population. This analysis (adjusted by COPD severity and asthma comorbidity) showed that patients who were black and had COPD were more likely to have an episode as a hospital inpatient, be it for any cause (OR 1.5 [95% CI = 1.0 to 2.1]) or for respiratory-specific causes (OR 2.3 [95% CI = 1.1 to 4.6]).

**DISCUSSION**

Summary

The results of this study demonstrate clear differences in COPD prevalence, smoking status, severity, management, and hospital admissions by ethnicity. COPD among the white population was recorded in younger groups and had a greater prevalence at all ages in comparison with other ethnic groups in east London. COPD in the white population was equally distributed by sex, while for black and South Asian groups both smoking rates and COPD were more common among males. Among those with COPD, smoking rates remained much higher in the white population, in comparison with other groups. Both South Asian and black populations reported less severe breathlessness symptoms than their white counterparts; this could suggest that they have a milder COPD or that symptoms are better tolerated or managed in comparison with the white population. These findings showed that, for COPD drug prescribing, there was a trend for the black population to be on less medication, regardless of symptoms or severity; in contrast, South Asians were largely on a similar range of medications — apart from lower rates of LAMA. South Asian patients were also more likely to have influenza and pneumococcal vaccinations than other groups.

Among the Tower Hamlets population, black patients were more likely to be admitted to hospital for any cause, and for COPD-related admissions than white or South Asian groups. This may suggest these
groups have more frequent and severe exacerbations, and may be linked to the observed lower rates of prescribed medications.

**Strengths and limitations**

The high levels of self-reported ethnicity in this study population enable robust estimation of the prevalence of COPD by ethnicity. However, the low overall prevalence suggests that case identification in general practice is suboptimal. Improving the rate of detection of COPD towards the estimated prevalence of 3% identified by the British Lung Foundation is now an important task in primary care.17

Study limitations include a reliance on routine clinical data entry during consultations, which may have introduced data of varying quality. Spirometry values recorded in the previous 2 years were available for 61% of cases, so there is the possibility of diagnostic inaccuracy within the cohort. A lack of routinely recorded data on pack years means there is a dearth of detail on the variation in smoking intensity between groups. In addition, differences could exist in the perception or reporting behaviour by ethnicity that might explain the differences in symptoms that were noted.

**Comparison with existing literature**

Variation in prevalence by ethnicity has been observed both in the UK1 and the US.10 But, as far as the authors are aware, this is the first attempt to examine COPD management by ethnicity in UK primary care. Simpson et al found higher rates of smoking continuation associated with social deprivation in England, but did not include analysis by ethnicity.11 It is possible that these differences may reflect variation in smoking habits between ethnic groups; information on pack years was not available, which may be an important variable in influencing smoking cessation.

Although there is a lack of research into differences in COPD management according to ethnicity, there is evidence from studies on other chronic diseases, particularly cardiovascular disease, which demonstrate similar patterns in the use of prescribed medicines; that is, higher rates among South Asians and lower rates among black populations.21–23 The reasons for these differences are uncertain. Given the equitable access to care in the UK NHS, differential registration and access to primary and secondary care services are unlikely to play a role; differential take-up of medicines due to cost is also unlikely in this population as the majority are of retirement age and, hence, eligible for free prescriptions. It is possible that differences in health beliefs and the understanding of COPD between groups may influence medication use and approaches to self-management. Studies in asthma have identified lower levels of confidence in self-managing acute attacks among South Asians, which contributes to higher rates of hospital admission than white groups.25

**Implications for practice**

Improving the primary care management of patients with COPD is important both for optimal symptom management and to reduce the rate of exacerbation and the need for hospital admission. Drug management in this cohort was suboptimal, with surprisingly low rates of SABA prescribing. In contrast, there were high rates of IC prescribing, either alone or in combination with a LABA, in spite of the fact that only 30.6% of cases had an FEV1% of <50. The most recent NICE guidance8 supports the use of combined LABAs and ICs in moderate but symptomatic or exacerbating COPD, but this does come with the risk of increased rates of pneumonia.

Recorded asthma comorbidity was high in this cohort and particularly so among the South Asian population (data available from the author on request). It is unclear whether this was due to diagnostic uncertainty or genuine differences in comorbidity. Referral rates to pulmonary rehabilitation at 18.5% remain low, particularly among minority ethnic groups, although these rates are an improvement on the national 5% referral rates quoted in 2006.26

Equitable access and utilisation of healthcare remains a priority for the Department of Health, and the recent Marmot review emphasises the importance of identifying and monitoring inequalities as a first stage to intervention.27 Routine audit data, such as the QOF in the UK, does not include ethnicity as a measure, which restricts its use as a tool for commissioning in multiethnic areas and as a framework for monitoring inequalities by ethnicity in the management of chronic disease. The findings of this study will be relevant to other areas in the UK that have large minority ethnic populations, but differences in the management of COPD by ethnicity need to be explored further, and addressed, in order to reduce outcomes inequalities. Further work is needed to understand the differences in prescribing rates by ethnicity, and how this relates to the severity of symptoms and perceptions of the disease in different groups.
REFERENCES


