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A comparison of medication adherence/persistence for asthma and chronic obstructive pulmonary disease in the United Kingdom

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Abstract

AIM: To describe and compare adherence and persistence with inhaled therapies in patients with asthma or COPD in the United Kingdom (UK).

METHODS: A retrospective prescribing database cohort was obtained from 44 general practitioner surgeries in NHS Forth Valley Scotland. Patients with physician-diagnosed asthma or COPD who received inhaled therapy between January 2008 and December 2009 were included. Four classes of inhaled therapy were assessed: inhaled corticosteroids; long-acting beta-agonists; combination therapy inhalers and; long-acting muscarinic antagonists. Adherence was calculated using the medication possession ratio (MPR) and persistence was determined using Kaplan-Meier survival analysis for the time to discontinuation (TTD) over one year. Two step-wise logistic regressions were performed to assess the contribution of diagnosis to adherence/persistence.

RESULTS: 12,923 patients were included in the analysis: 10,177 patients with asthma and 2,746 patients with COPD. 24.8% of medication episodes for asthma and 45.0% of medication episodes for COPD were classified as having an adequate medication supply (MPR of 80-120%). The overall median TTD was 90 days (IQR: 50-184 days) for patients with asthma and 115 days (58-258 days, comparison p<0.001) for patients with COPD. Patients with COPD were found to be more likely to achieve an MPR of at least 80% (OR: 1.32, 95% CI: 1.20-1.46), but had a similar likelihood of persistence at one year to patients with asthma.

CONCLUSION: Adherence and persistence with respiratory therapies in the UK is relatively low. There is suggestion that patients with COPD may display more adherent behaviours than patients with asthma.
What is already known about this topic:

- Patients with respiratory disease are particularly likely to have poor adherence to medicines due to complex medication regimens and difficulty using inhaler devices.
- It is unknown whether patients with asthma and COPD have different medication use behaviours.

What this article adds:

- Adherence for maintenance inhaler therapies in respiratory disease is low, but higher among patients with COPD compared to patients with asthma.
- Less than 20% of patients in this UK cohort with respiratory disease remained persistent with inhaled therapy at one year.
Main text

INTRODUCTION:

Medication adherence refers to the degree to which a patient's medication-taking behaviour coincides with agreed medical advice, and largely has supplanted the term 'compliance,' which infers a clinician-directed therapy decision without patient input.[1] Medication persistence examines whether a patient follows the recommendation of continuing treatment for a prescribed length of time.[2] According to the World Health Organization (WHO), approximately half of patients across the world are non-adherent with treatment prescribed for chronic disease, leading to decreased treatment effectiveness, decreased patient quality of life, and increased healthcare costs.[1] As clinical research develops new therapies for patients with chronic disease it is important to identify and optimise medication taking behaviours.[3-4]

Treatment non-adherence in respiratory disease is particularly common. For patients with asthma, reported non-adherence rates range from 30-70%[5] and are thought to contribute significantly to the prevalence of severe refractory disease.[6] Most studies quantifying adherence in asthma have focused on children although adults are of equal concern as non-adherence occurs across all demographic categories.[7] In patients with chronic obstructive pulmonary disease (COPD), complex medication regimens and multiple comorbidities result in adherence rates of approximately 50%.[8] A lack of perceived benefit led to 30% of patients with COPD intentionally discontinuing their therapy in one analysis.[9] For both conditions, rates of non-
adherence reported in clinical trials are likely to be underestimated due to the type of patients selected for trials and the influence of the trial environment itself.

Respiratory disease presents a unique challenge for patient adherence/persistence. In addition to the common barriers that all patients with chronic disease face, a number of device-related factors complicate the situation, including dislike of inhaled formulations, improper inhaler technique, and need for additional equipment such as spacers.[10] Of additional importance is how patients perceive their disease in terms of severity and awareness,[11] which for respiratory disease – with a variable symptom time course in asthma, and misunderstanding of disease symptoms as ‘normal’ in COPD – is an important contributor.

Almost all available respiratory medications are utilised in the treatment of both asthma and COPD. Accordingly, some studies have analysed adherence jointly without regard to diagnosis,[12-13] despite the different pathology, symptoms and management of each condition. To our knowledge, no single analysis has compared the differences in respiratory medication utilisation between patients with asthma or COPD, and accordingly, the aim of this study was to describe and compare adherence and persistence with inhaled therapies for patients with physician-diagnosed asthma and COPD in the United Kingdom (UK).

METHODS:

Patients

The present study is a retrospective database cohort analysis of prescribing data from National Health Services (NHS) Forth Valley in Scotland. Data were provided
by the NHS Forth Valley Airways Managed Clinical Network in coordination with the E-PRS clinical recording tool program (Campbell Software Solutions\textsuperscript{\textregistered}, Irvine, UK). E-PRS was designed as a computer-integrated clinical audit program to improve clinical care at practice level (described elsewhere\cite{14}) but in short, collectively formed a database of patients with physician-diagnosed asthma and/or COPD treated in 44 general practitioner (GP) surgeries from 2007 to 2009. The reference health board covers a geographic population of nearly 300,000 patients, while the database contains information on approximately 22,000 distinct patients with physician-diagnosed asthma/COPD. The departmental ethics group at the University of Strathclyde determined no formal ethics review was needed for use of the dataset.

Patients (children and adults) included in the analysis had (1) a GP-recorded diagnosis of asthma or COPD, and (2) at least one prescription issued for a qualifying inhaled maintenance medication between January 1, 2008, and December 31, 2009 (the study period), including inhaled corticosteroids (ICS), long-acting beta agonists (LABA), ICS/LABA combination therapy inhalers, or long-acting muscarinic antagonists (LAMA). Patients were assessed separately by diagnosis, and patients with a recorded dual-diagnosis of asthma and COPD were excluded from the analysis. Demographic characteristics for each group were assessed, including age, sex, maintenance medications received, and the proportion of patients who were prescribed reliever medications (short-acting beta agonists [SABA] and oral corticosteroids [OCS]) during the study period.

\textit{Adherence}
Medication adherence was assessed by calculating the medication possession ratio (MPR) for each class of maintenance therapy received. Similar to other analyses,[12,15-16] MPR was calculated by summing the days of medication supply provided and dividing by the total time treated:

\[
\text{MPR} = \frac{\text{total days of medication supply}}{\text{days between first and last fills}} \times 100\%
\]

Medication supply was calculated according to the posology and inhaler size (e.g. 60-dose inhaler with prescriptive instructions of 1 inhalation twice daily = 30 days supply) for each prescription. A sequence of at least two prescriptions during the two-year study period was required to calculate an MPR. The calculation was specific to therapeutic class, but not to dose or individual pharmacological agent. Thus, a patient receiving sequential treatment with two different ICS inhalers (e.g. fluticasone switched to beclometasone) or having a change in the dose mid-therapy would have a single MPR calculation. However, a single patient may have multiple MPRs if they received medications from more than one therapeutic class, such as treatment with an ICS changed to treatment with a combination therapy inhaler. After individual MPRs were calculated for each patient and medication, they were aggregated by therapeutic class: ICS, combination therapy inhalers, LABA (for patients with asthma or COPD), and LAMA (only for patients with COPD). The use of LAMA was not quantified for patients with asthma due to a lack of treatment recommendation in prevailing clinical guidelines. The proportion of patients having achieved an adequate medication supply, defined as an MPR between 80 and 120%,[17] was also determined; MPRs less than 80% or over 120% were defined as
undersupply or oversupply, respectively. The amount of SABA (in doses/day)
prescribed during each MPR was also quantified. This was achieved by summing the
number and size of SABA inhalers to determine the number of total SABA doses
prescribed, and then dividing by the number of days over which the maintenance
inhaler was prescribed (the denominator of the MPR calculation).

Persistence
Medication persistence was evaluated using a refill-sequence model and defined as
the difference in time between the first prescribing of a medication during the study
period and either the last prescribing, or an non-permissible gap in therapy
(whichever occurred first).[18-19] A patient was considered to have an unacceptable
gap if they failed to receive a prescription within 30 days after their previous
medication supply was due to run out. Patients were also classified by whether they
were on new or established therapy during the study period; patients were
considered to be on 'new' therapy if they had no history of being prescribed an agent
in the specified therapeutic class in the six months prior to the start of the study
period. At least one prescription for a maintenance medication during the first year of
the two-year study period was required to enable accurate calculation of a one-year
persistence rate. Median time to discontinuation (TTD) for each patient by
therapeutic class was calculated using a Kaplan-Meier survival analysis, with a time
censor utilised at one year post-medication initiation.

Statistical analysis
Two binary logistic regression models were utilised to separately assess predictors
of adherence and persistence. For adherence, the outcome of interest was
achievement of an MPR of at least 80% (including both adequate supply and oversupply); this binary measure was chosen over a continuous measure since oversupply does not confer any demonstrable therapeutic benefit over adequate supply. For persistence, a post hoc consideration of the Kaplan-Meier results suggested a non-proportionality of hazards over time; therefore, logistic regression (with an outcome of interest of persistence past 100 days of therapy) was utilised over a Cox regression analysis. Patient- and treatment-related variables of interest (age, sex, diagnosis, therapeutic class of medication) were assessed first in univariate fashion; SABA and OCS utilisation (doses/day and receipt [yes/no], respectively) during the study period were also entered in adherence model to include a measure of disease control, and classification of treatment as new/established therapy was utilised in the persistence model. Significant variables from the univariate analysis were filtered into the final forward stepwise multivariate models, with α set at 0.05 for entry and 0.10 for removal. Results were reported with odds ratios (OR) and 95% confidence intervals (CI).

Minitab® 16 statistical software (Minitab Ltd., Coventry, UK) was used for analysis. A Mann-Whitney test was used to compare differences between continuous measures (median age, median doses/day of SABA), while chi-squared tests were used to assess categorical differences (demographics, classification of medication supply). A Wilcoxon test was used to test for differences in persistence as assessed by the Kaplan-Meier analysis. A Bonferroni correction was applied to minimise error resulting from multiple comparisons when appropriate.

RESULTS:
Patient characteristics

A total of 12,923 patients were included in the analysis, 10,177 patients with asthma and 2,746 patients with COPD (Table 1). Patients with asthma were more likely to be female, and the median age of patients was predictably younger and more variable than those with COPD. More patients with asthma (8873; 87.2%) than with COPD (2746; 65.4%) were treated with a single therapeutic class of medication during the study period (p<0.001). Patients with either disease were prescribed SABA in similar proportions, although the use of OCS was greater among those with COPD. A total of 17,354 episodes of maintenance medication use were assessed: 11,618 for asthma patients and 5,736 for COPD patients.

Medication adherence

Overall 24.8% (95% confidence interval [CI]: 24.0-25.6%) of medication episodes for asthma and 45.0% (95% CI: 43.7-46.3%) of medication episodes for COPD were classified as an adequate supply (p<0.001 for comparison). The proportion of patients with an adequate MPR also varied according to therapeutic class (Table 2). Adequate supply was highest for LAMA inhalers in patients with COPD at 52.2%; patients with COPD had higher incidence of adequate supply and lower incidence of undersupply for all three therapeutic classes available for comparison. Undersupply accounted for approximately one-third to one-half of all episodes, with all therapies for patients with asthma having an undersupply rate of greater than 50%. Oversupply of medication was particularly common for prescribing of ICS in both diseases, at 27.0% for asthma and 34.8% for COPD.
In the logistic regression for adherence, both male sex and increasing age were associated with higher odds of achieving an MPR of at least 80% (Table 3). Both a diagnosis of COPD and treatment with LAMA were associated with higher odds, although these effects were softened in the multivariate analysis with inclusion of other variables. The number of doses/day of SABA prescribed during the study period increased alongside MPR, with each additional dose/day correlating to an 11% increase in the odds of achieving an adequate MPR; receiving a prescription for OCS was also associated with an increased odds of an adequate MPR, although this effect failed to meet significance for inclusion in the multivariate model.

**Medication persistence**

For asthma, the overall median TTD was 90 days (IQR: 50-184 days) and for COPD, the overall median TTD was 115 days (IQR: 58-258, comparison p<0.001); the percentage of patients persisting at one year was 11% and 16% for asthma and COPD, respectively (Figure 1). Persistence by therapeutic class was greater for patients with COPD than asthma for both LABA (96 vs. 61 days, p=0.008) and combination inhalers (116 vs. 85 days, p<0.001), but similar for ICS (101 vs. 100 days). The highest persistence overall was seen for LAMA inhalers in COPD, with a median TTD of 123 days and 18% of patients persisting after one year.

In the logistic regression for persistence, male sex and age were associated with higher odds of persisting past 100 days of therapy (Table 4). Therapy with a LAMA or ICS produced the similarly highest odds among therapeutic classes, but diagnosis
failed to meet significance for inclusion in the final model. Newly initiated therapy was associated with 19% lower odds of persisting past 100 days, compared to patients on already established therapy.

DISCUSSION:

The current analysis demonstrates a relatively low rate of adherence/persistence for inhaled therapies for the treatment of respiratory disease, and reveals that among use of the same medications, patients with asthma and COPD display different medication use behaviour.

Less than one-quarter of patients with asthma and less than one-half of patients with COPD were classified as having an adequate medication supply. Other studies using administrative dispensing databases have demonstrated similar rates to those found in this prescribing analysis.[12-13,15-16,20-21] However, to the best of our knowledge, no single analysis has demonstrated enhanced adherence among patients with COPD for the same inhaled therapies within an adjusted analysis. There is evidence that patients with COPD display more adherent behaviours as they age,[22] and similarly our analysis found that age over 40 years is an independent contributor to having an adequate medication supply. However, older patients also face unique barriers that complicate medication adherence and may not be captured with prescribing data, including cognitive decline, lack of dexterity and polypharmacy.[23] Patients with COPD may also experience more consistent and severe disease symptomology, and may be inclined to be more adherent than patients with asthma. Although the inclusion of patients prescribed more than
reliever therapy would presume that the population studied had a chronic form of asthma, the natural episodic time course of patient symptoms in asthma may have resulted in sporadic treatment periods interlaced with drug-free intervals, rendering a lower level of overall adherence.

Adherence to prescribed therapy has been shown to have an association with disease outcomes in respiratory disease, resulting in lower rates of emergency room visits and hospitalisations.[24-26] Other measures of disease control, such as exacerbation rates, may be less sensitive and require high levels of adherence to provide noticeable improvement.[27] Our multivariate analysis found that patients with higher level prescribing of SABA reliever therapy were more likely to have higher level prescribing of their maintenance therapy, a relationship that has been detected previously.[15,28] It may be expected that a high use of reliever therapy should correspond with poor adherence to maintenance therapy; accordingly, it has been suggested previously that symptomatic patients may have a weaker sense of control and ownership of their illness, resulting in lower motivation and overall medication adherence.[29] However, the direction of causation in the relationship between reliever and maintenance therapy is unclear, as patients who are symptomatic as defined by increased use of reliever therapy may be inclined to develop better adherence to their maintenance therapy.[1] The perceived necessity of a medication by the patient has also been shown to correlate with adherence[29] and the relative severity of the disease consequences[11,30] may further complicate the situation.
More than 80% of patients failed to remain on prescribed therapy at 12 months post-initiation, not unlike an analysis of Canadian claims data that found a range of 47 – 93% for non-persistence at one year among inhaled therapies.[31] Medication persistence calculations are sensitive to how stringent the definition is for an allowable medication gap. The current analysis set this threshold at 30 days; a post hoc sensitivity analysis extending this gap to 60 days increased the overall median TTD to 114 days (20% persisting at one year) for asthma and 152 days (22% persisting at one year) for COPD, respectively (data not shown). Despite these results, the enhanced effect of COPD diagnosis seen in the regression for adherence failed to replicate in the regression for persistence. Medication persistence is not only a measure of long-term adherence, but also of treatment stability, as the TTD may be influenced by patients with therapy changes, such as for patients with asthma ‘stepping up’ from ICS therapy alone to ICS/LABA combination inhaler therapy. The percentage of patients treated with more than one class of maintenance therapy was significantly higher among patients with COPD than with asthma and likely softened the effect of both COPD diagnosis and treatment with LAMA in the multivariate regression for persistence – both variables that were significant on a univariate basis, and significant within the adherence regression. The shape of persistence curves and the further lessened persistence among patients with newly initiated therapy suggests that the overall low persistence is primarily influenced by the large drop-off of patients early in therapy, often after the first couple of prescriptions. Accordingly, initiation of therapy appears to be an appropriate target for promoting good medicine use in patients with respiratory disease. Although no single set of interventions has been successful in consistently
improving adherence, [3] our results would suggest that research should focus efforts during the first three months of therapy.

Among the inhaled therapies assessed, LAMAs were associated with the best adherence/persistence, which may be a function of the increased convenience of once-daily dosing.[32] Medication adherence is known to decrease as dosing frequency increases,[33] and all other therapies in the present analysis have a standard twice-daily dosing regimen. Therapy with an ICS had the lowest percentage of patients (with either disease) identified as having an adequate level of medication supply (MPR 80-120%), but was also found to have a comparatively high rate of adherence/persistence in the regression analyses. For adherence, this was likely influenced by the inclusion of oversupply in the binary outcome, but for persistence, the reasons are less clear. There may have been some influence of inhaler size on the results, as most available combination therapy inhalers and LABAs are packaged in 60-dose or 120-dose units, largely corresponding to a 30-day supply per inhaler. However, the most commonly utilised ICSs in this analysis are supplied in 200-dose units, which may correspond to as much as a 100-day supply for a single inhaler. This may have led to higher odds of persistence for ICS within our calculations, as a single prescribing event provides a longer medication supply. An additional point to consider is the effect that ICS dose titration may have on adherence and persistence. It is possible that clinicians may initially prescribe a higher dose of ICS (or a higher number of puffs from an inhaler) which is then scaled back based on the patient’s symptoms; this may explain some variation in the results.
There are several limitations with the current study. All database studies estimate adherence/persistence as it is not always feasible to obtain serum drug concentrations or individually assess patients in large numbers. Access to medication is used as a proxy for adherence in this study and thus we are unable to discern whether the patient actually takes the medication or if he/she uses an inhaler device correctly; therefore, by definition the results are ceiling estimates.

Furthermore, the current study used prescribing data obtained from the GP. Prescriptions are generally written for an initial supply, and a patient only continues therapy by obtaining a new prescription, either by visiting the GP themselves, or more commonly through a 'managed repeat' service where the pharmacist communicates with the GP for further supply. While the first prescription is initiated by the GP, there is a great degree of patient-initiated action required for follow-up supply, which may influence the large-drop off seen early in therapy. Similarly, hospitalisations, particularly for patients with COPD may lead to changes or disruptions in therapy that effect the calculations. Lastly, the regression models provide an indication of the variables when considered together, but indicate (via the c-statistic) that other variables that were unable to be accounted for within this analysis likely have an effect on the outcome; however, this modest fit is a common finding among adherence studies.[34] In particular, in a prescribing database, we were unable to correlate the results of the regression with clinical outcomes, such as hospitalisations and overall disease control. However, we attempted to include a proxy for symptoms by quantifying SABA and OCS utilisation among patients.

In conclusion, adherence and persistence with respiratory therapies in this UK cohort was relatively low. There is indication that patients with COPD may display more
adherent behaviours than their counterparts with asthma. Efforts to increase medication persistence should likely be targeted to focus follow-up efforts during the first months of therapy.

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AUTHOR CONTRIBUTIONS:

JRC and ABM came up with the concept and designed the study. JRC, BFJ, ABM and DTS analysed the study data. ACB, FW and MR interpreted the results. JRC drafted the manuscript, and all authors critically reviewed and approved the manuscript. ACB is the guarantor.
REFERENCES:

1. World Health Organization (WHO). Adherence to long-term therapies: evidence for action. 2003. Available at:
   http://www.who.int/chp/knowledge/publications/adherence_full_report.pdf
   [accessed 12 Feb 2013]


34. Zeber JE, Manias E, Williams AF, Hutchins D, Udezi WA, Roberts CS, Peterson AM; ISPOR Medication Adherence Good Research Practices Working Group. A systematic literature review of psychosocial and behavioral...
Table 1: Patient characteristics

<table>
<thead>
<tr>
<th></th>
<th>Asthma (n=10,177)</th>
<th>COPD (n=2746)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>n (%) unless otherwise stated</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male sex</td>
<td>4317 (42.4)</td>
<td>1367 (49.8)‡</td>
</tr>
<tr>
<td>Median age, years (IQR)</td>
<td>41 (22 – 57)</td>
<td>70 (63 – 77)‡</td>
</tr>
<tr>
<td>&lt; 20 years</td>
<td>2204 (21.7)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>20-39 years</td>
<td>2561 (25.2)</td>
<td>7 (0.3)</td>
</tr>
<tr>
<td>40-59 years</td>
<td>3235 (31.8)</td>
<td>418 (15.2)</td>
</tr>
<tr>
<td>60-80 years</td>
<td>1900 (18.7)</td>
<td>1853 (67.5)</td>
</tr>
<tr>
<td>≥ 80 years</td>
<td>277 (2.7)</td>
<td>468 (17.0)</td>
</tr>
<tr>
<td><strong>Therapies received</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ICS</td>
<td>6520 (64.1)</td>
<td>682 (24.8)‡</td>
</tr>
<tr>
<td>CMB</td>
<td>4087 (40.2)</td>
<td>1504 (54.8)‡</td>
</tr>
<tr>
<td>LABA</td>
<td>874 (8.6)</td>
<td>274 (10.0)†</td>
</tr>
<tr>
<td>LAMA</td>
<td>N/A</td>
<td>1736 (63.2)</td>
</tr>
<tr>
<td>Received SABA</td>
<td>9456 (92.9)</td>
<td>2534 (92.3)</td>
</tr>
<tr>
<td>Received OCS</td>
<td>2426 (23.8)</td>
<td>1179 (42.9)‡</td>
</tr>
</tbody>
</table>

CMB: combination therapy inhaler; COPD: chronic obstructive pulmonary disease; ICS: inhaled corticosteroid; IQR: interquartile range; n: number; LABA: long-acting beta-agonist; LAMA: long-acting muscarinic antagonist; OCS: oral corticosteroids; SABA: short-acting beta agonists

† indicates p<0.05 for comparison with asthma group
‡ indicates p<0.001 for comparison with asthma group
Table 2: Medication supply as assessed by MPR, by therapeutic class and diagnosis

<table>
<thead>
<tr>
<th>n (%)</th>
<th>ICS ‡</th>
<th>LABA ‡</th>
<th>CMB ‡</th>
<th>LAMA</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Asthma</td>
<td>COPD</td>
<td>Asthma</td>
<td>COPD</td>
</tr>
<tr>
<td>Undersupply</td>
<td>3156 (50.4)</td>
<td>250 (34.2)</td>
<td>466 (57.6)</td>
<td>163 (44.9)</td>
</tr>
<tr>
<td>Adequate</td>
<td>1414 (22.6)</td>
<td>227 (31.0)</td>
<td>220 (27.2)</td>
<td>122 (33.6)</td>
</tr>
<tr>
<td>Oversupply</td>
<td>1692 (27.0)</td>
<td>255 (34.8)</td>
<td>123 (15.2)</td>
<td>78 (21.5)</td>
</tr>
</tbody>
</table>

CMB: combination therapy inhaler; COPD: chronic obstructive pulmonary disease; ICS: inhaled corticosteroid; n: number; LABA: long-acting beta-agonist; LAMA: long-acting muscarinic antagonist
‡ indicates p<0.001 for comparison within therapeutic class
Table 3: Results of logistic regression for adherence

<table>
<thead>
<tr>
<th>Variable</th>
<th>Crude OR (95% CI) †</th>
<th>Adjusted OR (95% CI) ‡§</th>
<th>Adjusted OR p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>1 (reference)</td>
<td>1 (reference)</td>
<td>0.013</td>
</tr>
<tr>
<td>Male</td>
<td>1.15 (1.08 – 1.23)</td>
<td>1.09 (1.02 – 1.16)</td>
<td></td>
</tr>
<tr>
<td>Age</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0-19 years</td>
<td>1 (reference)</td>
<td>1 (reference)</td>
<td></td>
</tr>
<tr>
<td>20-39 years</td>
<td>1.03 (0.93 – 1.15)</td>
<td>0.94 (0.84 – 1.06)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>40-59 years</td>
<td>1.27 (1.16 – 1.41)</td>
<td>1.17 (1.06 – 1.30)</td>
<td></td>
</tr>
<tr>
<td>60-79 years</td>
<td>1.85 (1.68 – 2.03)</td>
<td>1.51 (1.35 – 1.69)</td>
<td></td>
</tr>
<tr>
<td>80+ years</td>
<td>2.04 (1.75 – 2.37)</td>
<td>1.60 (1.35 – 1.89)</td>
<td></td>
</tr>
<tr>
<td>Diagnosis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Asthma</td>
<td>1 (reference)</td>
<td>1 (reference)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>COPD</td>
<td>1.99 (1.86 – 2.12)</td>
<td>1.32 (1.20 – 1.46)</td>
<td></td>
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<tr>
<td>Therapeutic class</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ICS</td>
<td>1 (reference)</td>
<td>1 (reference)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>LABA</td>
<td>0.83 (0.73 – 0.94)</td>
<td>0.68 (0.59 – 0.77)</td>
<td></td>
</tr>
<tr>
<td>CMB</td>
<td>1.02 (0.96 – 1.09)</td>
<td>0.87 (0.81 – 0.94)</td>
<td></td>
</tr>
<tr>
<td>LAMA</td>
<td>2.19 (1.98 – 2.42)</td>
<td>1.33 (1.17 – 1.52)</td>
<td></td>
</tr>
<tr>
<td>SABA (doses/day)</td>
<td>1.11 (1.10 – 1.12)</td>
<td>1.11 (1.10 – 1.12)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>OCS</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>1 (reference)</td>
<td>N/A</td>
<td>0.824</td>
</tr>
<tr>
<td>Yes</td>
<td>1.24 (1.16 – 1.31)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

CI: confidence interval; CMB: combination therapy inhaler; COPD: chronic obstructive pulmonary disease; ICS: inhaled corticosteroid; LABA: long-acting beta-agonist; LAMA: long-acting muscarinic antagonist; N/A: not applicable; OCS: oral corticosteroids; OR: odds ratio; SABA: short-acting beta agonist

† univariate analysis with MPR ≥ 80% utilised as outcome
‡ multivariate analysis adjusted by sex, age, diagnosis, therapeutic class and SABA utilisation, with MPR ≥ 80% utilised as outcome
§ model fit assessed by c-statistic: 0.68 (95% CI: 0.67-0.69)
Table 4: Results of logistic regression for persistence

<table>
<thead>
<tr>
<th>Variable</th>
<th>Crude OR (95% CI) †</th>
<th>Adjusted OR (95% CI) ‡§</th>
<th>Adjusted OR p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>1 (reference)</td>
<td>1 (reference)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Male</td>
<td>1.14 (1.07 – 1.21)</td>
<td>1.14 (1.07 – 1.21)</td>
<td></td>
</tr>
<tr>
<td>Age</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0-19 years</td>
<td>1 (reference)</td>
<td>1 (reference)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>20-39 years</td>
<td>0.91 (0.82 – 1.01)</td>
<td>0.97 (0.87 – 1.08)</td>
<td></td>
</tr>
<tr>
<td>40-59 years</td>
<td>1.09 (0.99 – 1.21)</td>
<td>1.17 (1.06 – 1.29)</td>
<td></td>
</tr>
<tr>
<td>60-79 years</td>
<td>1.48 (1.35 – 1.63)</td>
<td>1.58 (1.43 – 1.75)</td>
<td></td>
</tr>
<tr>
<td>80+ years</td>
<td>1.53 (1.31 – 1.79)</td>
<td>1.61 (1.37 – 1.90)</td>
<td></td>
</tr>
<tr>
<td>Diagnosis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Asthma</td>
<td>1 (reference)</td>
<td>N/A</td>
<td>0.444</td>
</tr>
<tr>
<td>COPD</td>
<td>1.31 (1.22 – 1.41)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Therapeutic class</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ICS</td>
<td>1 (reference)</td>
<td>1 (reference)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>LABA</td>
<td>0.66 (0.58 – 0.74)</td>
<td>0.60 (0.53 – 0.68)</td>
<td></td>
</tr>
<tr>
<td>CMB</td>
<td>0.89 (0.83 – 0.95)</td>
<td>0.81 (0.76 – 0.88)</td>
<td></td>
</tr>
<tr>
<td>LAMA</td>
<td>1.23 (1.11 – 1.37)</td>
<td>0.94 (0.84 – 1.06)</td>
<td></td>
</tr>
<tr>
<td>Type of therapy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Established</td>
<td>1 (reference)</td>
<td>1 (reference)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>New</td>
<td>0.78 (0.72 – 0.84)</td>
<td>0.81 (0.75 – 0.87)</td>
<td></td>
</tr>
</tbody>
</table>

CI: confidence interval; CMB: combination therapy inhaler; COPD: chronic obstructive pulmonary disease; ICS: inhaled corticosteroid; LABA: long-acting beta-agonist; LAMA: long-acting muscarinic antagonist; N/A: not applicable; OR: odds ratio

† univariate analysis with persistence > 100 days utilised as outcome
‡ multivariate analysis adjusted by sex, age, therapeutic class and type of therapy, with persistence > 100 days utilised as outcome
§ model fit assessed by c-statistic: 0.57 (95% CI: 0.56-0.58)
Kaplan-Meier survival curves for 1-year medication persistence for asthma and COPD patients overall