∂ Open Access Full Text Article

ORIGINAL RESEARCH

Economic impact of delaying initiation with multiple-inhaler maintenance triple therapy in Spanish patients with chronic obstructive pulmonary disease

This article was published in the following Dove Press journal: International Journal of Chronic Obstructive Pulmonary Disease

Antoni Sicras Mainar¹ Alicia Huerta² Ruth Navarro Artieda³ Eduard Monsó^{4,5} Sarah H Landis⁶ Afisi S Ismaila^{7,8}

¹Scientific Direction, Health Economics and Outcomes Research (HEOR) Department, Real Life Data, Madrid, Spain; ²Market Access Department, GlaxoSmithKline SA, Madrid, Spain; ³Medical Documentation Department, Hospital Germans Trias I Pujol, Badalona, Spain; ⁴Pulmonology Service, Hospital Parc Taulí, Barcelona, Spain; ⁵CIBERES -Ciber De Enfermedades Respiratorias, Madrid, Spain; ⁶Real World Evidence and Epidemiology Department, GlaxoSmithKline, Uxbridge, UK; ⁷Value Evidence and Outcomes Department, GlaxoSmithKline, Collegeville, PA, USA; ⁸Department of Health Research Methods, Evidence and Impact, McMaster University, Hamilton, ON, Canada

Purpose: Guidelines recommend the use of triple therapy with an inhaled corticosteroid (ICS), a long-acting β_2 agonist (LABA) and a long-acting muscarinic antagonist (LAMA) to reduce the risk of future exacerbations in symptomatic COPD patients with a history of exacerbations. This study aimed to estimate COPD-related healthcare resource use and costs, and subsequent exacerbation rates, for patients initiating multiple-inhaler triple therapy (MITT) early (\leq 30 days) versus late (31–180 days) following an exacerbation, in a real-world clinical setting.

Patients and methods: This was an observational, longitudinal, retrospective study using electronic medical records from the Spanish database of the Red de Investigación en Servicios Sanitarios Foundation. Patients \geq 40 years old with a confirmed COPD diagnosis who were newly prescribed MITT up to 180 days after an exacerbation between January 2013 and December 2015 were included. Patients were followed from the date of MITT initiation for up to 12 months to assess COPD-related health care resource use (routine and emergency visits, hospitalizations, pharmacologic treatment), exacerbation rate, and costs (€2017); these endpoints were compared between early versus late groups.

Results: The study included 1280 patients who met selection criteria: mean age 73 years, 78% male, and 41% had severe/very severe lung function impairment. The proportion of patients initiating MITT early versus late was 61.6% versus 38.4%, respectively. There were no statistically significant differences in baseline characteristics between groups. During follow-up, health care resource consumption was lower in the early versus late group, especially primary care and ED visits, leading to lower total costs (€1861 versus €1935; P<0.05). In the follow-up period, 28.0% of the patients in the early group experienced ≥1 exacerbation versus 36.4% in the late group (P=0.002), with an exacerbation rate of 0.5 versus 0.6 per person per year (P=0.022), respectively.

Conclusion: Initiating MITT early (\leq 30 days after an exacerbation) may reduce health care costs and exacerbation rate compared with late MITT initiation.

Keywords: COPD, triple therapy, costs, exacerbations, inhaled corticosteroid

Correspondence: Alicia Huerta Market Access Department, GlaxoSmithKline SA, C/Severo Ochoa 2, Madrid 28760, Spain Tel +34 91 807 0821 Fax +34 9 187 0500 Email donha_ali@hotmail.com



Introduction

Chronic obstructive pulmonary disease (COPD) is a common, preventable, and treatable disease characterized by persistent respiratory symptoms and airflow limitation, usually caused by exposure to noxious particles or gases.¹ It is one of the leading causes of mortality and morbidity worldwide.

International Journal of Chronic Obstructive Pulmonary Disease 2019:14 2121–2129 2121 © 2019 Sicras Mainar et al. This work is published and licensed by Dove Medical Press Limited. The full terms of this license are available at https://www.dovepress.com/ the work you hereby accept the Terms. Non-commercial uses of the work are permitted without any further permission from Dove Medical Press Limited, provided the work is properly attributed. For permission for commercial use of this work, please see paragraphs 4.2 and 5 of our Terms (https://www.dovepress.com/terms.php). Exacerbations are a key clinical feature of COPD and are recognized as one of the main drivers of the burden of the disease. They are defined as an acute worsening of respiratory symptoms that result in additional therapy² and can cause increased decline of lung function, impairment of health status, and increased mortality.^{3–5} On average, patients can experience 1–4 exacerbations per year, though this is highly variable, with history of previous exacerbations being the strongest predictor of a patient's future exacerbation frequency.⁶ Exacerbations are also associated with substantial economic burden; those requiring hospital admission are responsible for an important part of COPD-attributable costs. Indeed, they may account for up to 50–75% of the total costs of the disease.⁷

While corticosteroids and antibiotics are crucial for the acute treatment of exacerbations, appropriate pharmacologic maintenance treatment should also be initiated for prevention of future exacerbations. Specifically, treatment with the triple therapy combination of an inhaled corticosteroid (ICS) with two bronchodilators, a long-acting β_2 agonist (LABA) and a long-acting muscarinic antagonist (LAMA), has been shown to reduce exacerbation rate versus dual therapy with ICS and a bronchodilator.^{8,9} As such, the Global Strategy for the Diagnosis, Management, and Prevention of COPD (GOLD) 2019 report recommends escalating to triple therapy to reduce the risk of future exacerbations in COPD patients with persistent symptoms who experience exacerbations on dual LAMA/LABA or ICS/LABA therapy.¹

Evidence exploring whether the timing of triple therapy initiation influences the risk of future exacerbations (and therefore the overall costs associated with the disease) is scarce, especially in a real-world setting, with only one recent article addressing this topic.¹⁰ The aim of this study was to estimate the economic and clinical (exacerbation rate) impact of initiating multiple-inhaler triple therapy (MITT) early (\leq 30 days) versus late (31–180 days) following a COPD-related exacerbation in a routine clinical practice setting in Spain.

Materials and methods

Study design and patient population

This was an observational, longitudinal, retrospective database study based on the review of electronic medical records (EMRs) linked from the patient database of the Red de Investigación en Servicios Sanitarios (RedISS) Foundation, a research-based organization that uses

EMRs stored via OMIAP software (in use at many health care centers in Spain), as well as other complementary databases. More specifically, the RedISS database contains data on patient demographic characteristics, medical conditions recorded using the International Statistical Classification of Diseases and Related Health Problems codes (ninth revision; ICD-9-CM), clinical variables, medical services dispensed (date and location of medical services, type and level of service, including emergency departments), hospitalizations (date of admission and discharge, primary diagnoses, medical services and interventions), days of sick leave, as well as prescribed and dispensed medication (date of service, name of treatment, dosage strength and amount dispensed, type of prescription and duration of treatment). The data are representative of the Spanish population.

The confidentiality of the records established by the Spanish Organic Data Protection Law (15/1999 of 13 December) was respected by dissociation of the data. The study was classified as a post-authorization study-other design (EPA-OD) by the Spanish Agency for Medicines and Medical Devices and subsequently approved by the Ethics Committee of Universitat Internacional de Catalunya.

The patient identification period was 1 January 2013 to 31 December 2015. During that period, the index date was defined as the date of inpatient discharge from the first COPD exacerbation-related hospitalization or hospital emergency department (ED) visit followed by initiation of MITT in a 180-day window (Figure 1). For the purposes of this study, MITT was defined as an overlapping prescription of an ICS/LABA fixed-dose combination plus a LAMA single agent, or an overlapping prescription of a LAMA/LABA fixed-dose combination plus an ICS single agent, or an overlapping prescription of ICS, LAMA, and LABA single agents.

The timing of MITT prescription from the patient identification index date was used to define two study groups: an early cohort (prescription \leq 30 days from the index date) and a late cohort (prescription 31–180 days from the index date). Patients were followed up for study outcomes, including exacerbations and health care resource use, starting from the date of MITT initiation for up to 12 months.

The study population included patients \geq 40 years old with a COPD diagnosis identified by ICD-9-CM codes 491.x and 492.x and confirmed with registry of a forced expiratory volume in 1 second (FEV₁)/forced vital capacity (FVC) ratio

2122

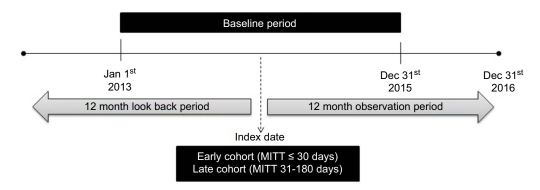


Figure I Study design. Abbreviation: MITT, multiple-inhaler triple therapy.

<0.70, who were prescribed an MITT treatment up to 180 days after an exacerbation that required hospitalization or an ED visit, and who had available medical records during the 12month follow-up period (except in case of death). Patients were excluded from the study if they had a prescription record of a MITT treatment in the 12 months prior to the index date or if they initiated MITT treatment >180 days after the index date.

Demographic and clinical variables

The following demographic and clinical variables were collected in order to describe the study sample: age (continuous and by group), gender, occupational status (active worker or retiree), body mass index (BMI), smoking status, lung function impairment (last FEV₁ registry), time from diagnosis, presence of comorbidities such as hypertension, type 2 diabetes mellitus, dyslipidemia, obesity, smoking habit, alcoholism, organ failure (heart, liver or kidneys), ischemic cardiomyopathy, cerebrovascular accident, dementia, psychosis, neurological disorders (Parkinson's disease, epilepsy or multiple sclerosis), depressive syndrome, malignant neoplasms, and substance abuse. In addition, the Charlson comorbidity index¹¹ and the number of comorbidities were collected as an estimation of the seriousness of the patient's condition.

Outcomes

Exacerbation rate

Exacerbations events were collected from the MITT initiation date for up to 12-months' follow-up for both the early and late cohorts.

Two types of exacerbations were considered: a) severe: those that required hospital admission and b) moderate: those that required treatment with antibiotics or oral corticosteroids (criteria based on the health care resource use). Exacerbations were identified by diagnostic code (496.x) and by treatment (prescription for antibiotics and oral corticosteroids in the absence of another codified infectious event, such as tonsillitis or urine infection). Exacerbation episodes that included hospital admission were classified as severe exacerbations; exacerbations without hospital admission were classified as moderate exacerbations. The moderate/ severe exacerbation rate was calculated for each study group and expressed per person per year.

Health care utilization and costs

COPD-related health care resources, including visits to primary and secondary care, medical procedures, ED visits, hospitalizations and pharmacologic treatment were also collected from the MITT initiation date for up to 12months' follow-up.

Total COPD-related direct costs throughout the analysis period (12 months from MITT initiation) were quantified by multiplying the natural unit of the health care resource used by the unit cost of each resource. Unit costs were obtained from analytical accounting of the centers. In the case of medication, the price to public (PTP) plus the value-added tax (VAT) price at the moment of dispensation was used.¹² All costs were inflation adjusted to \notin 2017 (Table 1).

Treatment persistence

Treatment persistence (or duration in days) of MITT was calculated from index date until discontinuation date. The discontinuation date was the first of the following events: a) the end date of the 12-month follow-up period (continuous use through the whole period); b) change to another treatment different from MITT; and/or c) interrupt/withdraw of MITT treatment (≥ 60 days without renewing the treatment).

Table I Unit costs

Healthcare resources	Unit costs (€)
Medical Visits	
Medical visit to primary care	23.19
Medical visit to emergency department	117.53
Medical visit to secondary care ^a	67.50
Hospitalization (1 day)	320.90
Complementary tests	
Laboratory tests	22.30
Conventional radiology	18.50
Diagnostic/therapeutic tests	37.12
Spirometry	15.00
Computerized axial tomography	92.00
Magnetic resonance	154.00
Bronchodilator tests	56.00
Pharmacologic treatment	PTP+VAT

Notes: Values are expressed in \in 2017. ^aOnly pulmonologist and internal medicine were considered. Data from analitic accounting of centers included in the RedISS database.¹²

Abbreviations: PTP, price to public; VAT, value added tax.

Results were presented as the percentage of patients achieving the end of follow-up period treated with MITT.

Statistical analysis

A descriptive univariate statistical analysis was performed to characterize the study sample, and both the early and late cohorts. Continuous variables were described using mean and standard deviation, and median and interquartile range. Categorical variables were described using absolute and relative frequencies. Univariate comparison between groups was conducted using independent sample *t*-test and Wilcoxonsigned ranked test for continuous variables, and Fisher's exact test and chi-square test for categorical variables.

A comparison of resource use and its corresponding cost was performed according to the recommendations of Thompson and Barber¹³ using a general linear model (ANCOVA). The covariates used in the model were gender, age, time since diagnosis, Charlson index, and number of exacerbations in the follow-up period.

Additionally, the time to first exacerbation was estimated using Kaplan–Meier survival analysis. A cox proportional hazard model corrected for covariates (age, gender, comorbidities, time from diagnosis, lung function, previous treatments, history of exacerbations) was used to estimate differences in time-to-first exacerbation by study groups (HR: hazard ratio).

The software program used was SPSS version 23, with statistical significance established for P-values <0.05.

Results

From an initial selection of 209,557 subjects \geq 18 years old included in the database, 9,805 had a registered COPD diagnosis and 1,280 met the study criteria and were included in the analysis. In total, 76.2% of the patients had a COPD exacerbation that required an ED visit and 23.8% had a COPD exacerbation that required hospitalization. Of patients with an exacerbation event, 788 (61.6%) initiated MITT treatment early (\leq 30 days) following the exacerbation versus 492 (38.4%) who initiated late (31–180 days). Regarding MITT treatment, 61.3% of the patients were prescribed ICS/LABA plus LAMA, 35.2% LAMA/LABA plus ICS, and 3.5% ICS plus LAMA plus LABA.

Table 2 shows the baseline demographics and clinical characteristics of the study sample and study groups. The mean age was 73 years, 78% were men, and the mean Charlson index score was 1. Arterial hypertension (59%) and dyslipidemia (50%) were the most frequent comorbidities. Mean FEV₁ was 54% and 41% of the patients had severe/ very severe lung function impairment. Regarding treatment persistence to MITT, 68.6% of the patients treated with ICS/ LABA plus LAMA, 67.9% of the patients treated with LAMA/ LABA plus ICS, and 67.1% of the patients treated with ICS plus LAMA plus LABA were persistent to treatment. There were no statistically significant differences between groups.

During the 12-month follow-up period, health care resource use was lower in the early versus the late group, especially primary care visits (12.8 versus 15.8, P<0.001) and secondary care visits (1.6 versus 1.9, p<0.001) (Table 3). Of the patients included in the study, the total costs amounted to €2.4 million, of which 80% was from primary care (including pharmacologic costs) and 20% from secondary care. The main contributors of costs were pharmacologic treatment (57.8%), primary care visits (17.1%), and hospitalization (11.4%).

By group, total costs were lower in the early versus the late group ($(\in 1,823 \text{ versus } \in 1,959, P=0.041$). These differences were maintained after adjusting for covariates (Table 4). The adjusted average total cost per patient in the early group was $(\in 1,861 (\in 1,766-(\in 1,956) \text{ versus } \in 1,935 (\in 1,820-(\in 2,049) \text{ in the})$ late group (mean average difference of $(\in 74, P=0.046)$). By cost component, all costs were lower in the early group except for pharmacologic treatment. In the general linear model, cost was associated (P<0.001) with the number of exacerbations ($\beta=0.628$), severity of the disease ($\beta=0.124$), age ($\beta=0.117$), and FEV₁ ($\beta=-0.087$). The determinant coefficient of the model was 47.1%.

Table 2 Baseline demographics and clinical characteristics of the study sample and per study groups

Study group	Total N=1,280 (100%)	Early N=788 (61.6%)	Late N=492 (38.4%)	P
Socio-demographic characteristics				
Age (years), mean (SD, SE)	73.2 (10.4; 0.36)	73.3 (10.2; 0.38)	73.0 (10.7; 0.30)	0.579
40–64 years	20.5%	20.1%	21.3%	
65–74 years	30.5%	30.5%	30.7%	
≥75 years	48.9%	49.5%	48.0%	
Gender (male)	77.9%	78.9%	76.2%	0.255
Retiree status	85.2%	85.8%	84.1%	0.422
General comorbidity		•		
Charlson index score, mean (SD, SE)	0.9 (0.8; 0.01)	0.9 (0.9; 0.01)	0.9 (0.8; 0.01)	0.629
Number of diagnoses, mean (SD, SE)	3.8 (1.9; 0.07)	3.8 (1.9; 0.08)	3.7 (1.9; 0.05)	0.455
I	9%	8.8%	9.3%	
2	16.3%	15.1%	18.1%	
3	18.5%	19.4%	17.1%	
4	19.9%	17.9%	23.2%	
≥5	36.3%	38.8%	32.3%	
Associated comorbidities				
Hypertension	59.0%	59.4%	58.3%	0.708
Type 2 diabetes mellitus	32.7%	33.5%	31.5%	0.459
Dyslipidemia	50.2%	50.6%	49.4%	0.665
Obesity	35.0%	36.2%	33.1%	0.268
Active smoker	14.1%	14.1%	14.2%	0.976
Ischemic cardiomyopathy	19.2%	19.7%	18.5%	0.604
Cerebrovascular accident	13.3%	13.6%	12.8%	0.691
Cardiac insufficiency	18.0%	17.9%	18.0%	0.960
Renal insufficiency	12.6%	12.4%	12.8%	0.847
Depressive syndrome	23.6%	23.5%	23.8%	0.851
Malignant neoplasms	15.8%	15.1%	16.9%	0.399
Osteoporosis	15.7%	15.4%	16.3%	0.655
Clinical characteristics				I
Time since diagnosis (years), mean (SD, SE)	17.1 (3.9; 0.15)	17.2 (4.5; 0.13)	17.0 (2.9; 0.12)	0.404
BMI, kg/m ² , mean (SD, SE)	28.8 (5.2; 0.18)	28.9 (5.1; 0.20)	28.7 (5.2; 0.12)	0.573
FEVI, % (SD, SE)	54.1 (14.4; 0.51)	54.0 (14.3; 0.64)	54.4 (14.4; 0.40)	0.649
Moderate	58.7%	57.9%	60.0%	
Severe	38.8%	40.0%	37.0%	
Very severe	2.5%	2.2%	3.0%	
Exacerbation events in previous 12 months				
Moderate or severe	37.0%	35.4%	39.0%	0.194
Moderate	36.7%	34.8%	38.5%	0.147
Severe	12.8%	11.3%	14.2%	0.153
Treatment persistence				
ICS/LABA plus LAMA	68.6%	68.8%	67.3%	0.668
LAMA/LABA plus ICS	67.9%	67.2%	69.3%	0.728
ICS plus LAMA plus LABA	67.1%	67.5%	66.4%	0.831

Note: Values are expressed as percentage or mean (SD; SE).

Abbreviations: BMI, body mass index; FEV1, forced expiratory volume in one second; SD, standard deviation; SE, standard error; ICS, inhaled corticosteroid; LABA, longacting $\beta 2$ agonist; LAMA, long-acting muscarinic antagonist.

Table 3 Healthcare resource use during 12-month follow-up period per study group

Study group	Early N=788 (61.6%)	Late N=492 (38.4%)	Р
Primary care visits	12.8 (11.8; 0.42)	15.8 (13.8; 0.52)	<0.001
Laboratory tests	0.6 (0.9; 0.04)	0.8 (1.1; 0.05)	0.002
Chest radiology	0.3 (0.8; 0.02)	0.5 (0.8; 0.03)	<0.001
Complementary tests	1.7 (1.5; 0.05)	1.9 (1.5; 0.06)	0.020
Hospitalization	0.1 (0.3; 0.01)	0.1 (0.4; 0.01)	0.011
Secondary care visits	1.6 (1.6; 0.05)	1.9 (1.8; 0.08)	0.001
Emergency visits (hospital)	0.3 (0.8; 0.02)	0.5 (0.8; 0.03)	<0.001

Note: Values are expressed as mean (SD; SE).

Abbreviations: SD, standard deviation; SE, standard error.

Table 4 Health care costs per study group

Study group	Early N=788 (61.6%)	Late N=492 (38.4%)	Р
Uncorrected costs (€)			
Costs in primary care	1,505 (527; 18.7)	1,495 (563; 25.4)	0.742
Visits	294 (272; 9.6)	364 (318; 14.3)	<0.001
Laboratory tests	30 (42; 1.4)	37 (49; 1.2)	0.002
Conventional radiology	6 (14; 0.5)	10 (16; 0.6)	<0.001
Complementary tests	54 (45; 1.6)	60 (48; 2.1)	0.020
Pharmacologic treatment	1,122 (354; 9.5)	1,024 (364; 9.9)	<0.001
Costs in secondary care	317 (737; 26.2)	464 (899; 40.5)	0.002
Hospitalization	173 (644; 22.9)	277 (798; 35.9)	0.011
Visits	104 (104; 3.7)	125 (117; 5.2)	0.001
Emergency visits	41 (95; 3.3)	61 (96; 4.3)	<0.001
Total uncorrected costs	1,823 (1,078; 38.4)	1,959 (1,265; 57.0)	0.041
Corrected costs ^a (€)	·		
Primary care costs	1,500	1,471	0.367
95% CI	1,459–1,542	1,420–1,521	
Secondary care costs	361	464	0.042
95% CI	292-428	381–546	
Total corrected costs	1,861	1,935	0.046
95% CI	1,766–1,956	1,820–2,049	

Notes: Uncorrected costs are expressed as mean (SD; SE). ^aANCOVA model, the contrasts are based on the comparison by linear independent pairs between the estimated marginal means. Variables are gender, age, Charlson index score, time from diagnosis, and number of exacerbations. **Abbreviations:** CI, confidence interval; SD, standard deviation; SE, standard error.

During the follow-up period, 31.3% of the patients experienced at least one moderate/severe exacerbation. Fewer patients (28.0%) in the early cohort suffered ≥ 1 exacerbation versus the late cohort (36.4%, *P*=0.002), with a lower mean rate of exacerbations (0.5 versus 0.6, *P*=0.022). Considering only patients who suffered an exacerbation during the follow-up period, the mean rate of exacerbations was 1.6 in the early group versus 1.8 in the late group (*P*=0.028). These differences were also observed when exacerbations were separated into moderate and severe (Table 5). The median time from MITT initiation to first exacerbation was 130 days in the early cohort versus 80 days in the late cohort (P=0.001; Figure 2); the HR for the late cohort compared to the early cohort was 0.74 (95% CI: 0.62–0.89).

Discussion

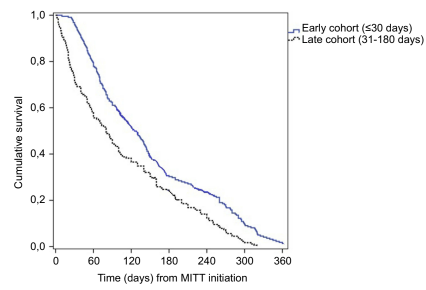
This study evaluated the impact of early versus late initiation with MITT after an exacerbation that required hospitalization or a hospital ED visit and showed that initiation

Table 5 Exacerbations during 12-month follow-up period per study group

Study group	Early N=788 (61.6%)	Late N=492 (38.4%)	
Patients with exacerbations	28.0%	36.4%	0.002
Exacerbation rate per person per year	0.5 (1.0; 0.01)	0.6 (1.0; 0.01)	0.022
0	72.0%	63.6%	
I	14.3%	16.3%	
2	6.5%	14.0%	
>3	7.2%	6.1%	
Patients with moderate exacerbations	27.4%	36.0%	0.005
Moderate exacerbation rate per person per year	0.4 (0.8; 0.02)	0.5 (0.8; 0.02)	0.037
0	72.6%	64.0%	
I	17.4%	26.2%	
2	5.8%	5.7%	
≥3	4.2%	4.1%	
Patients with severe exacerbations	27.4%	36.0%	0.005
Severe exacerbation rate per person per year	9.0%	14.4%	0.004
0	0.1 (0.3; 0.02)	0.1 (0.4; 0.02)	0.011
I	92.5%	87.8%	
2	6.0%	10.2%	
Time to first exacerbation, days			
Mean (SD)	148.2 (96.3)	107.1 (90.5)	<0.00
Median (IQR)	130 (66–217)	80 (28–173)	

Note: Values are expressed as percentages or mean (SD; SE).

Abbreviations: IQR, interquartile range; SD, standard deviation; SE, standard error.



Kaplan-meier curve - log-rank procedure (mantel-cox): chi-squared=20.643, P=0.001

Figure 2 Time to first exacerbation per study group.

within 30 days of the exacerbation resulted in decreased health care resource use, lower exacerbation rates, and lower COPD-related direct costs compared with late initiation of MITT. The average difference in costs between early and late patients in this study could be considered modest (\notin 74 per patient), mainly due to the

fact that the costs avoided in the early group are counterbalanced by the extra pharmacologic cost.

Triple therapy is becoming increasingly important in clinical practice. A retrospective cohort study of newly diagnosed COPD patients performed in the UK showed that 23% were initially treated with MITT and 24% of the remaining patients were escalated onto triple therapy within the following 24 months.¹⁴ In Spain, an epidemiologic study performed in the primary care setting showed that 12% of newly diagnosed COPD patients initiated triple therapy.¹⁵ This percentage increased to 49% in patients treated in outpatient respiratory clinics.¹⁶

GOLD guidelines recommend the use of triple therapy in COPD patients with persistent symptoms who experience exacerbations on dual LAMA/LABA or ICS/LABA therapy. Understanding the pathway and drivers of treatment escalation to triple therapy is key. In this respect, a recent study showed that COPD exacerbations were the most significant factor for escalation.¹⁷ In another study,¹⁸ results showed that exacerbations had only a minor influence on treatment augmentation and that it was driven more by the need for rescue medication and assessment of dyspnea. In our study of patients with COPD experiencing an exacerbation event, it can be considered that exacerbations were the driver of MITT initiation and, in these cases, initiation sooner after the exacerbation had beneficial effects versus later MITT initiation.

Similar to the findings reported here, a study performed in the USA¹⁰ showed that the total costs for COPD patients who delayed triple therapy initiation were 18.7% higher than the costs for patients who initiated triple therapy promptly. Other studies have also examined the potential benefits of initiating an ICS-containing regimen along with bronchodilators. One study showed that early treatment with ICS/LABA reduced severe exacerbations and COPDrelated costs,¹⁹ supporting the idea that reducing any delay in treatment intervention may have beneficial effects.

In this study, triple therapy was administered via two or three inhalers, and there may have been different frequencies of administration. Single-inhaler triple therapy was recently launched in Spain, after the period assessed in the present study, with the potential of improving adherence, reducing complexity, and thus improving patient outcomes.

The mean annual cost per patient was $\notin 1875$ (result not shown). This result is in line with previous publications that estimated the cost of COPD management in Spain. De Miguel-Diaz et al²⁰ found that the annual cost of a COPD patient in primary care was $\notin 1920$, with hospitalizations

and drug costs being the larger components of costs. In our study, the main drivers of costs were also pharmacologic treatment and primary care visits; costs were associated with the number of exacerbations, disease severity, and age. The impact of exacerbations on costs has been previously studied. Another study, also performed in Spain, showed that patients with exacerbations had 2.3 times more resource consumption than patients who had not suffered any exacerbations.²¹

The findings of this study should be considered within the context of its limitations. First, there could have been a cohort misclassification if a patient classified as "late" had initiated MITT treatment during hospitalization, had enough medication to last 1 month and thus did not fill a prescription for MITT in the first 30 days. The impact of this misclassification would have reduced the treatment effect of early initiation. Second, as it is a retrospective database study, there could have been misclassification or underascertainment of COPD or exacerbations. This, and poor registration of health care resources consumed outside the centers that were part of the database, could have led to an underestimation of costs. However, this limitation would have impacted both groups equally. Lastly, although the analysis was adjusted by covariates, there could be some residual confounding that was not considered.

Conclusion

Initiation of MITT treatment \leq 30 days after a COPD exacerbation-related hospitalization or ED visit may reduce health care costs and exacerbation rates compared with late MITT initiation in a routine clinical practice setting.

Acknowledgment

The abstract of this paper was presented at the ISPOR Europe Congress 2018 as a poster presentation with interim findings. The poster's abstract was published in Value in Health 2018; 21, sup 3:S407: https://tools.ispor.org/ScientificPresentationsDatabase/Presentation/85607? pdfid=58276.

Disclosure

ASI is employe of, and hold stocks in, GlaxoSmithKline plc. ASI is also unpaid part-time faculty at McMaster University, ON, Canada. SHL and AH were employees of GlaxoSmithKline plc. at the time of the study. ASM and EM received fees for their participation in the study. The authors report no other conflicts of interest in this work.

References

- Global strategy for the diagnosis, management and prevention of chronic obstructive pulmonary disease. 2019 Report. Global Initiative for Chronic Obstructive Lung Disease. Available from: http://www.goldcopd.org. Accessed August 06, 2019.
- Wedzicha JA, Seemungal TA. COPD exacerbations: defining their cause and prevention. *Lancet*. 2007;370(9589):786–796. doi:10.1016/ S0140-6736(07)61382-8
- Donaldson GC, Seemungal TA, Bhowmik A, Wedzicha JA. Relationship between exacerbation frequency and lung function decline in chronic obstructive pulmonary disease. *Thorax*. 2002;57 (10):847–852. doi:10.1136/thorax.57.10.847
- Miravitlles M, Ferrer M, Pont A, et al. Effect of exacerbations on quality of life in patients with chronic obstructive pulmonary disease: a 2 year follow-up study. *Thorax.* 2004;59(5):387–395. doi:10.1136/ thx.2003.008730
- Soler-Cataluna JJ, Martínez-García MM, Román-Sánchez P, Salcedo E, Navarro M, Ochando R. Severe acute exacerbations and mortality in patients with chronic obstructive pulmonary disease. *Thorax*. 2005;60:925–931. doi:10.1136/thx.2005.040527
- Hurst JR, Vestbo J, Anzueto A, et al. Susceptibility to exacerbation in chronic obstructive pulmonary disease. N Engl J Med. 2010;363 (12):1128–1138. doi:10.1056/NEJMoa0909883
- Celli BR, MacNee W. Standards for the diagnosis and treatment of patients with COPD. A summary of the ATS/ERS position paper. *Eur Respir J.* 2004;23:932–946.
- Lipson DA, Barnhart F, Brealey N, et al. Once-daily single-inhaler triple versus dual therapy in patients with COPD. N Engl J Med. 2018;378(18):1671–1680. doi:10.1056/NEJMoa1713901
- Papi A, Vestbo J, Fabbri L, et al. Extrafine inhaled triple therapy versus dual bronchodilator therapy in chronic obstructive pulmonary disease (TRIBUTE): a double-blind, parallel group, randomised controlled trial. *Lancet*. 2018;391(10125):1076–1084. doi:10.1016/ S0140-6736(18)30206-X
- Bogart M, Glassberg MB, Reisch T, Stanford RH. Impact of prompt versus delayed initiation of triple therapy post COPD exacerbation in a US-managed care setting. *Respir Med.* 2018;145:138–144. doi:10.1016/j.rmed.2018.10.013
- Charlson ME, Pompei P, Ales KL, Mackenzie CR. A new method of classifying prognostic comorbidity in longitudinal studies: development and validation. *J Chronic Dis.* 1987;40(5):373–383.

- Ministerio de Sanidad Servicios Sociales e Igualdad. Information about the products included in the pharmaceutical provision of the SNS (dispensable through pharmacies). 2018. Available from: http://www.msssi. gob.es/en/profesionales/nomenclator.do. Accessed August 06, 2019.
- Thompson SG, Barber JA. How should cost data in pragmatic randomised clinical trials be analysed? *BMJ*. 2000;1197–1200. doi:10.1136/bmj.320.7243.1197
- Wurst KE, Punekar YS, Shukla A. Treatment evolution after COPD diagnosis in the UK primary care setting. *PLoS One*. 2014;9(9): e105296. doi:10.1371/journal.pone.0105296
- Barrecheguren M, Monteagudo M, Ferrer J, et al. Treatment patterns in COPD patients newly diagnosed in primary care. A population-based study. *Respir Med.* 2016;111:47–53. doi:10.1016/j.rmed.2015.12.004
- Calle Rubio M, Alcázar Navarrete B, Soriano JB, et al. Clinical audit of COPD in outpatient respiratory clinics in Spain: the EPOCONSUL study. Int J Chron Obstruct Pulmon Dis. 2017;12:417–426. doi:10.2147/COPD.S124482
- Hurst JR, Dileen M, Morris K, Hills S, Emir B, Jones R. Factors influencing treatment escalation from long-acting muscarinic antagonist monotherapy to triple therapy in patients with COPD: a retrospective THIN-database analysis. *Int J Chron Obstruct Pulmon Dis.* 2018;13:781–792. doi:10.2147/COPD.S153655
- Landis SH, Wirst K, Le HV, Bonar K, Punekar YS. Can assessment of disease burden prior to changes in initial COPD maintenance treatment provide insight into remaining unmet needs? A retrospective database study in UK primary care. *COPD*. 2017;14(1):80–85. doi:10.1080/15412555.2016.1240159
- Akazawa M, Biddle AK, Stearns SC. Economic assessment of early initiation of inhaled corticosteroids in chronic obstructive pulmonary disease using propensity score matching. *Clin Ther.* 2008;30:Spec No:1003–1016. doi:10.1016/j.clinthera.2008.05.020
- 20. de Miguel Diez J, Carrasco Garrido P, García Carballo M, et al. Determinants and predictors of the cost of COPD in primary care: a Spanish perspective. *Int J Chron Obstruct Pulmon Dis.* 2008;3 (4):701–712.
- Sicras A, Huerta A, Navarro R, Ibáñez J. Use of resources and associated costs of chronic obstructive pulmonary disease exacerbations: a population based retrospective study. *Semergen*. 2014;40:189–197. doi:10.1016/j.semerg.2013.10.002

International Journal of Chronic Obstructive Pulmonary Disease

Dovepress

Publish your work in this journal

The International Journal of COPD is an international, peer-reviewed journal of therapeutics and pharmacology focusing on concise rapid reporting of clinical studies and reviews in COPD. Special focus is given to the pathophysiological processes underlying the disease, intervention programs, patient focused education, and self management

protocols. This journal is indexed on PubMed Central, MedLine and CAS. The manuscript management system is completely online and includes a very quick and fair peer-review system, which is all easy to use. Visit http://www.dovepress.com/testimonials.php to read real quotes from published authors.

Submit your manuscript here: https://www.dovepress.com/international-journal-of-chronic-obstructive-pulmonary-disease-journal-obstructive-pulmonary-disease-journal-obstructive-pu