Long-term outcomes and adverse events of therapy with inhaled corticosteroids, longacting beta-2-agonists and anticholinergic drugs in hospitalised patients with Chronic Obstructive Pulmonary Disease (COPD) - a cohort study based on health information systems in three Italian regions (OUTPUL)

First published: 27/10/2010 Last updated: 30/03/2015

Study Finalised

### Administrative details

#### **EU PAS number**

EUPAS1578

#### Study ID

9133

#### DARWIN EU® study

No

#### **Study countries**

Italy

#### **Study description**

Objectives: to measure long-term outcomes and adverse events of inhaled drugs in COPD patients, to compare effectiveness of the different drugs (monoand polytherapy) in terms of long-term survival or exacerbations, to compare incidence of side effects of inhaled therapy among users vs non-users. Design: patients discharged with COPD in 2006-07 will be enrolled from the Hospital Information Systems (HIS) and record linkage performed with mortality, hospital, emergency and drug claims data to define exposure (ICS, LABA, and anticholinergics), potential confounders, and to measure outcomes over 4 years. Outcomes: all-cause, respiratory and cardiovascular mortality, incidence of adverse events, and COPD exacerbations. Cox-proportional-hazard models will be applied to compare outcomes in users vs non-users of exposure to different drug groups (mono-/polytherapy). Sensitivity analyses will take into account different subgroup susceptibility or different definitions of chronic exposure.

#### **Study status**

Finalised

### Research institutions and networks

### Institutions

Department of Epidemiology of the Regional Health Service - Lazio

Italy
First published: 23/03/2010
Last updated: 22/06/2018
Institution EU Institution/Body/Agency ENCePP partner

Emilia-Romagna Health and Social Agency (ASSR Emilia-Romagna)

**ENCePP** partner

ltaly

First published: 23/04/2010

Last updated: 18/12/2017

Institution

ig( Laboratory/Research/Testing facility ig)

Epidemiologic Observatory of the Health Directorate of the Lombardy region, Milan, Italy, Respiratory Physiology Unit in the Columbus Hospital, Catholic University, Rome, Italy

## Contact details

Study institution contact Nera Agabiti u.kirchmayer@deplazio.it



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Primary lead investigator Nera Agabiti Primary lead investigator

### Study timelines

Date when funding contract was signed Planned: 01/06/2010 Actual: 16/06/2010

Study start date Planned: 15/09/2010 Actual: 15/09/2010

**Data analysis start date** Planned: 01/01/2012 Actual: 02/01/2012

**Date of interim report, if expected** Planned: 31/12/2011

Actual: 31/12/2011

Date of final study report Planned: 01/06/2013 Actual: 01/06/2014

## Sources of funding

• Other

### More details on funding

AIFA (Italian Medicines Agency), Participating centres

# Regulatory

Was the study required by a regulatory body?

Yes

Methodological aspects

Study type

# Study type list

#### Study topic:

Human medicinal product Disease /health condition

#### Study type:

Non-interventional study

#### Scope of the study:

Assessment of risk minimisation measure implementation or effectiveness Effectiveness study (incl. comparative)

#### Data collection methods:

Secondary use of data

#### Main study objective:

To measure long-term outcomes and adverse events of inhaled drugs in the study cohort over a 4-year follow up period, to compare effectiveness of the different drugs (both "monotherapy" and "combined therapy") in terms of longterm survival or exacerbations, to compare the incidence of side effects of inhaled therapy (ICS, LABA and anticholinergics) among users versus non users.

### Study Design

#### Non-interventional study design

Cohort

### Study drug and medical condition

#### Anatomical Therapeutic Chemical (ATC) code

(R03) DRUGS FOR OBSTRUCTIVE AIRWAY DISEASES
DRUGS FOR OBSTRUCTIVE AIRWAY DISEASES
(H02AB) Glucocorticoids
Glucocorticoids
(J01) ANTIBACTERIALS FOR SYSTEMIC USE
ANTIBACTERIALS FOR SYSTEMIC USE

#### Medical condition to be studied

Chronic obstructive pulmonary disease

## Population studied

#### Short description of the study population

Patients aged > 45years and resident in the study areas with acute exacerbation of COPD during a two-year period (2006-07)

#### Age groups

Adults (46 to < 65 years) Adults (65 to < 75 years) Adults (75 to < 85 years) Adults (85 years and over)

#### Special population of interest

Other

#### Special population of interest, other

Chronic obstructive pulmonary disease (COPD) patients

#### Estimated number of subjects

40000

# Study design details

#### Outcomes

Mortality (All causes, respiratory causes, cardiovascular causes), COPD exacerbation defined as either a hospital admission/emergency visit for COPD or COPD-related causes or a prescription for an oral corticosteroids with/without systemic antibiotic, Adverse cardio- and cerebrovascular events, Pneumonia, Osteoporotic fractures

#### Data analysis plan

Cox proportional hazard models will be applied to compare outcomes in users vs non-users of exposure to different drug groups (mono-/polytherapy). Sensitivity analyses will take into account different subgroup susceptibility or different definitions of chronic exposure.

### Documents

#### **Study publications**

Di Martino M1, Agabiti N, Bauleo L, Kirchmayer U, Cascini S, Pistelli R, Colame...

### Data management

## **ENCePP** Seal

#### This study has been awarded the ENCePP seal



### Data sources

#### Data sources (types)

Administrative healthcare records (e.g., claims) Drug dispensing/prescription data

# Use of a Common Data Model (CDM)

#### **CDM** mapping

No

# Data quality specifications

#### **Check conformance**

Unknown

#### **Check completeness**

Unknown

#### **Check stability**

Unknown

#### **Check logical consistency**

Unknown

## Data characterisation

#### Data characterisation conducted

No