Evidence in real world for Trixeo® Aerosphere™ Initiation in COPD (ENARXI)

First published: 27/09/2023

Last updated: 23/04/2024





Administrative details

EU PAS number EUPAS106900	
Study ID 106901	
DARWIN EU® study	
Study countries France	

Study description

The main aim of the study is to describe the characteristics of COPD patients when Trixeo® Aerosphere m is initiated, based on data from the THIN® and Colibri databases. Secondary objectives are: - To describe from THIN®

database data the rate of appropriate initiation of Trixeo® Aerosphere™ according to: - The Transparency Commission (CT) (18); - SPLF 2021 (5) and GOLD 2021 (4) recommendations. - To describe from THIN® database data the distribution of blood eosinophil count in patients initiating Trixeo® Aerosphere™. - To describe the characteristics of COPD patients initiating treatment with Trixeo® Aerosphere™ using data from the Colibri-BPCO cohort to compare the results obtained with THIN® data and supplement them with patients seen mainly by hospital and private lung specialists and potentially more severe.

Study status

Ongoing

Research institutions and networks

Institutions



Contact details

Study institution contact

Gaetan Deslée gdeslee@chu-reims.fr

Study contact

gdeslee@chu-reims.fr

Primary lead investigator

Adrien Coriat

Primary lead investigator

Study timelines

Date when funding contract was signed

Actual: 01/07/2021

Study start date

Actual: 01/09/2021

Date of final study report

Planned: 31/12/2023

Sources of funding

• Pharmaceutical company and other private sector

More details on funding

AstraZeneca

Regulatory

Was the study required by a regulatory body?

Yes

Is the study required by a Risk Management Plan (RMP)?

Not applicable

Methodological aspects

Study type

Study type list

Study type:

Non-interventional study

Scope of the study:

Drug utilisation

Main study objective:

The main aim of the study is to describe the characteristics of COPD patients when Trixeo® Aerosphere m is initiated, based on data from the THIN® database.

Study Design

Non-interventional study design

Cohort

Study drug and medical condition

Medical condition to be studied

Chronic obstructive pulmonary disease

Population studied

Age groups

Adults (18 to < 46 years)

Adults (46 to < 65 years)

Adults (65 to < 75 years)

Adults (75 to < 85 years)

Adults (85 years and over)

Estimated number of subjects

600

Study design details

Data analysis plan

No hypotheses will be tested and no assumptions will be made in this analysis. Continuous variables will be described using mean and standard deviation for parametric variables, median and interquartile range for non-parametric variables. Ordinal, binomial and categorical variables will be described using numbers and proportions.

Data management

Data sources

Data source(s) THIN® (The Health Improvement Network®)		
- Time (The fields)	improvement wetworks;	
Data source(s),	other	
Colibri		
Data sources (ty	pes)	
Electronic healthc	are records (EHR)	
Use of a Co	mmon Data Model (CDM)	
CDM mapping		
No		
Data qualit	y specifications	
Charle as of summer		
Check conforma	nce	
Unknown		
Check complete	ness	
Unknown		

Check stability

Unknown

Check logical consistency

Unknown

Data characterisation

Data characterisation conducted

No