Drug Use Investigation of ANORO ELLIPTA inhaler (200311)

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Administrative details

EU PAS number

EUPAS9868

Study ID

48235

DARWIN EU® study

No

Study countries

Japan

Study description

This investigation will be conducted to collect and evaluate information regarding the safety and efficacy of ANORO ELLIPTA under the actual postmarketing use conditions of the product.

Study status

Finalised

Research institutions and networks

Institutions

GlaxoSmithKline (GSK)

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Institution

Iwamizawa Municipal General Hospital Iwamizawa, Japan

Contact details

Study institution contact GSK Clinical Disclosure Advisor GSK Clinical Disclosure Advisor Pharma.CDR@gsk.com

Study contact

Pharma.CDR@gsk.com

Primary lead investigator

GSK Clinical Disclosure Advisor GSK Clinical Disclosure

Advisor

Primary lead investigator

Study timelines

Date when funding contract was signed Planned: 15/03/2013 Actual: 15/03/2013

Study start date Planned: 20/02/2015 Actual: 20/02/2015

Data analysis start date Actual: 03/07/2019

Date of final study report Planned: 31/12/2019 Actual: 31/10/2019

Sources of funding

• Pharmaceutical company and other private sector

More details on funding

GlaxoSmithKline

Study protocol

gsk-200311-protocol-redact.pdf(205.63 KB)

Regulatory

Was the study required by a regulatory body?

Yes

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

Non-EU RMP only

Methodological aspects

Study type

Study type list

Study topic: Human medicinal product

Study type:

Non-interventional study

Scope of the study:

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

Data collection methods:

Main study objective:

This investigation will be conducted to collect and evaluate information regarding the safety and efficacy of ANORO ELLIPTA under the actual postmarketing use conditions of the product.

Study Design

Non-interventional study design

Other

Non-interventional study design, other

Non interventional, Observational Post Marketing Surveillance under actual drug use condition.

Study drug and medical condition

Name of medicine

ANORO

Population studied

Short description of the study population

This investigation will be conducted in patients who are first prescribed Anoro for the approved indication of the product, "Relief of symptoms of obstructive airway disorder due to COPD (chronic bronchitis and emphysema) (in the case that long-acting inhaled anticholinergic and long-acting inhaled beta2-agonist combination is required)".

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)

Special population of interest

Hepatic impaired

Pregnant women

Renal impaired

Estimated number of subjects

2000

Study design details

Outcomes

Information regarding the safety and efficacy of ANORO ELLIPTA under the actual post-marketing use conditions of the product.

Data analysis plan

Items related to patient dispositionPatient demographic and baseline characteristicsItems related to safetyItems related to efficacy

Documents

Data management

Data sources

Data sources (types)

Other

Data sources (types), other Prospective patient-based data collection

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