# A cross-sectional study to evaluate the effectiveness of XALKORI Therapeutic Management Guide among physician prescribing XALKORI in Europe

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## Administrative details

EU PAS number	
EUPAS7389	
Study ID	
18665	
DARWIN EU® study	
No	
Study countries	
Austria	
Belgium	
Denmark	

France
Germany
☐ Ireland
Italy
Netherlands
Sweden
United Kingdom
Study status
Finalised
Research institutions and networks
Institutions
ICON Commercialisation & Outcomes



Multiple centres: 60 centers are involved in the study

## Contact details

## **Study institution contact**

Huang Kui kui.a.huang@pfizer.com

Study contact

kui.a.huang@pfizer.com

## **Primary lead investigator**

Huang Kui

**Primary lead investigator** 

## Study timelines

## Date when funding contract was signed

Planned: 28/07/2014 Actual: 18/08/2014

#### Study start date

Planned: 22/09/2014 Actual: 30/09/2014

### **Date of final study report**

Planned: 31/03/2017 Actual: 30/03/2017

# Sources of funding

• Pharmaceutical company and other private sector

## More details on funding

Pfizer Inc

## Study protocol

A8081049\_PROTOCOL\_Crizotinib Physician Survey\_25March2014.pdf(958.16 KB)

A8081049\_PROTOCOL AMENDMENT 2\_Crizotinib Physician Survey\_30 March 2015 Clean\_register.pdf(346.74 KB)

# Regulatory

Was the study required by a regulatory body?

Yes

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

EU RMP category 3 (required)

# Methodological aspects

Study type

Study type list

**Study topic:** 

Human medicinal product

#### Study type:

Non-interventional study

#### Scope of the study:

Effectiveness study (incl. comparative)

Other

#### If 'other', further details on the scope of the study

Evaluate the effectiveness of physician educational materials and assess whether physicians would provide patients with patient educational materials.

#### **Data collection methods:**

Primary data collection

#### Main study objective:

The overall objective of this study is to evaluate the effectiveness of the XALKORI TMG and PIB implemented to mitigate the risks of visual disorders, QTc prolongation, bradycardia, hepatotoxicity, neutropenia and leukopenia, and ILD/pneumonitis in 6 countries including Belgium, Denmark, France, Germany, Italy, and the Netherlands in the European Union (EU).

## Study Design

## Non-interventional study design

Cross-sectional

# Study drug and medical condition

Study drug International non-proprietary name (INN) or common name CRIZOTINIB

# Population studied

#### Short description of the study population

Physicians who have prescribed XALKORI per SmPC at least once within 12 months prior to taking the survey.

#### 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**

150

## Study design details

#### Data analysis plan

The survey population will include all physicians who are screened and eligible for this study. All statistical analyses in this study will be descriptive. All variables collected are categorical. Frequencies and percentages, with 95% Cls where appropriate, will be presented. Country specific analyses will be presented. Additional exploratory analyses and sensitivity analyses may be conducted.

## **Documents**

### Study results

Crizotinib A8081049 NI Study Report.pdf(6.54 MB)

## Data management

## **ENCePP Seal**

The use of the ENCePP Seal has been discontinued since February 2025. The ENCePP Seal fields are retained in the display mode for transparency but are no longer maintained.

## Data sources

**Data sources (types)** 

Other

Data sources (types), other

Physician survey

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**

Unknown