# Medical Records Review to Describe the Patterns of KRAS Testing and Vectibix Use in Europe (20101120)

First published: 11/02/2014

Last updated: 02/07/2024





# Administrative details

PURI
https://redirect.ema.europa.eu/resource/20214
EU PAS number
EU PAS IIuliibei
EUPAS5681
Study ID
20214
DARWIN EU® study
No
Study countries
Study countries
Belgium

Czechia
Denmark
France
Germany
Italy
Netherlands
Spain
Sweden
Study description
This non-interventional cross-sectional medical record review study will assess
the prevalence of KRAS testing and the impact of the KRAS test result on
patterns of Vectibix use in patients with metastatic colorectal cancer (mCRC)
treated with Vectibix in selected European countries over 3 rounds. As the
optimal use of Vectibix also requires accurate KRAS mutation testing, this study
will also assess data from the laboratory that performed the KRAS test. The
study will also monitor changes in the pattern of Vectibix treatment between
the different rounds of the study.
Study status
Finalised
Research institutions and networks
Institutions
Amgen
United States
United States

First published: 01/02/2024

Last updated: 21/02/2024



# Contact details

# **Study institution contact**

Global Development Leader Amgen, Inc

Study contact

medinfo@amgen.com

# **Primary lead investigator**

Global Development Leader Amgen, Inc

**Primary lead investigator** 

# Study timelines

Date when funding contract was signed

Actual: 15/09/2011

## Study start date

Actual: 19/09/2012

### Data analysis start date

Actual: 12/09/2013

# Date of interim report, if expected

Planned: 05/05/2014 Actual: 30/06/2014

# Date of final study report

Planned: 26/11/2015 Actual: 23/11/2015

# Sources of funding

• Pharmaceutical company and other private sector

# More details on funding

**Amgen** 

# Study protocol

Vectibix 20101120 Protocol 11Feb14.pdf(312.98 KB)

# Regulatory

Was the study required by a regulatory body?

Yes

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

EU RMP category 1 (imposed as condition of marketing authorisation)

# Methodological aspects

# Study type

Study type list

### **Study topic:**

Disease /health condition

Human medicinal product

Other

# Study topic, other:

Diagnostic procedure - KRAS testing

# Study type:

Non-interventional study

# Scope of the study:

Drug utilisation

Other

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

Describe patterns of KRAS biomarker testing prior to initiating treatment

### **Data collection methods:**

Secondary use of data

## Main study objective:

This non-interventional cross-sectional medical record review study will assess the prevalence of KRAS testing and the impact of the KRAS test result on patterns of Vectibix use in patients with metastatic colorectal cancer (mCRC) treated with Vectibix over 3 rounds. The study will also monitor changes in the pattern of Vectibix treatment between the different rounds of the study.

# Study Design

# Non-interventional study design

Cross-sectional

# Study drug and medical condition

### Name of medicine

**VECTIBIX** 

### Medical condition to be studied

Colorectal cancer metastatic

# Population studied

### Short description of the study population

Patients with metastatic colorectal cancer (mCRC) treated with Vectibix following the changes of label about the risk of Vectibix use in mCRC patients with mutant KRAS tumors in selected European countries.

Patients who had received Vectibix for the treatment of mCRC during the 6month period prior to the time when medical records were obtained, not have been in any experimental clinical trial at time of receiving Vectibix, and not have participated in this study, in an earlier round were included.

### 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

Other

# Special population of interest, other

Colorectal cancer patients

### **Estimated number of subjects**

450

# Study design details

### **Outcomes**

The prevalence of KRAS testing and impact of the KRAS test results on patterns of Vectibix use in patients with metastatic colorectal cancer (mCRC) treated with Vectibix Timeframe: 3 Years, The proportion of oncologists who participate in the study. The proportion of oncologists who conduct a KRAS test prior to initiating Vectibix treatment. The proportion of labs that are part of an European Society of Pathology Quality Assurance scheme. The proportion of labs that use a validated KRAS testing method. To assess the change in pattern of Vectibix administation over the 3 rounds.

### Data analysis plan

Analysis will be conducted on 3 levels - patient level, oncologist level and laboratory level. Descriptive statistics will be calculated at each level.

# **Documents**

# Study results

Observational Research Study Report 20101120 - Abstract.pdf(35.72 KB)

# Data management

# Data sources

Data sources (types) Other	
Data sources (types), other Cross-sectional medical chart review	
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