Evaluate the Real-World Safety Outcomes and Clinical Efficacy of Ponatinib and Other Tyrosine Kinase Inhibitors among Chronic Myeloid Leukemia Patients

First published: 15/02/2023 Last updated: 02/07/2024



## Administrative details

#### **EU PAS number**

EUPAS50308

#### **Study ID**

50309

#### DARWIN EU® study

No

#### **Study countries**

United States

### **Study description**

The aims of this study are to learn out about treatment information (including amongst others treatment patterns, safety, development of a participant's condition) ponatinib, bosutinib, imatinib, dasatinib and nilotinib using already available data. No new data will be collected from participants as part of this study and no study medicines will be provided in this study.

#### Study status

Finalised

## Research institutions and networks

## Institutions

### Takeda

First published: 01/02/2024

Last updated: 01/02/2024

Institution

# Contact details

### Study institution contact

Study Contact Takeda TrialDisclosures@takeda.com

Study contact

TrialDisclosures@takeda.com

Primary lead investigator

Study Contact Takeda

Primary lead investigator

# Study timelines

**Date when funding contract was signed** Actual: 06/10/2021

Study start date Actual: 24/07/2020

Data analysis start date Actual: 24/07/2020

Date of interim report, if expected Actual: 22/12/2021

Date of final study report Actual: 30/11/2022

## Sources of funding

• Pharmaceutical company and other private sector

### More details on funding

Takeda

## Study protocol

Takeda\_Protocol\_CML\_Humedica\_24JULY2020\_redacted\_12 Jan 2024.pdf(4.79 MB)

# Regulatory

#### Was the study required by a regulatory body?

No

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

Not applicable

## Methodological aspects

# Study type

# Study type list

### Study topic:

Disease /health condition

#### Study type:

Non-interventional study

### Scope of the study:

Disease epidemiology Effectiveness study (incl. comparative)

#### Data collection methods:

Secondary use of data

#### Main study objective:

The main objective of this study is using Humedica Electronic Health Records (EHR) data from October 1, 2012—September 30, 2017 (or the most recent 5 years of data), STATinMED Research proposes to evaluate the real-world treatment patterns and clinical outcomes of ponatinib and other Tyrosine Kinase Inhibitors (TKIs) among patients with prior TKI uses among CP-CML participants.

# Study Design

### Non-interventional study design

Cohort

## Study drug and medical condition

#### Medical condition to be studied

Chronic myeloid leukaemia

## Population studied

#### Short description of the study population

The study population included patients with chronic myeloid leukemia received treatment with ponatinib and other tyrosine kinase inhibitors identified from Humedica EMR data between October 1, 2012 to September 30, 2017.

### 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

Patients with chronic myeloid leukemia

### Estimated number of subjects

1769

# Study design details

### Outcomes

Participants Categorized by Socio-demographic Variables, Clinical Characteristics of Disease,Comorbidities Severity,Concomitant Medication,Bone Marrow Stem Cell Transplant,Major Adverse Cardiac Event(MACE),Arterial Occlusive Event(AOE), and Venous Thrombotic Events(VTE),Quan-Charlson Comorbidity Index Score,Number of Previous Treatments of TKI Drugs,Duration From Last TKI Run-out to Index Date. Participants With BCR-ABL,Bone Marrow Testing, Disease Severity as per Medstat Disease Staging Clinical Criteria Version 5.21, Treatment-Free Gap of the Index Treatment,Treatment Patterns Based on Duration of Index Treatment,Mean Starting Daily Dose and Average Daily Dose,Number of Participants With CML on Concomitant Medication.

#### Data analysis plan

All variables will first be analyzed descriptively and compared across ponatinib, bosutinib, and other TKIs (imatinib, dasatinib, or nilotinib) for patients with prior TKI use. Numbers and percentages will be provided for dichotomous and polychotomous variables. Means and standard deviations will be provided for continuous variables. For dichotomous and polychotomous variables, p-values will be calculated according to the chi-square test, for continuous variables, ttests will be used to calculate p-values. Ponatinib will be considered as the reference group for the comparisons.

## Documents

#### **Study report**

GOR-2017-102256-clinical-study-report-redact.pdf(856.19 KB)

### Data management

### Data sources

#### Data source(s), other

Humedica EMR

#### Data sources (types)

Administrative healthcare records (e.g., claims) Electronic healthcare records (EHR)

## 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