

### Randomized, Open-Label, Phase II, Multicenter, Multi-Country Study to Evaluate Safety and Efficacy of Dasatinib 50 mg in First -Line Treatment of Early Chronic Phase Chronic Myeloid Leukemia

07/08/2025 17:03:29

Main		

Primary registry identifying number

LBCTR2019010169

MOH registration number

Study registered at the country of origin

Type of registration

Prospective

Date of registration in national regulatory

05/11/2018

**Primary sponsor** 

Hikma Pharmaceuticals

Date of registration in primary registry

10/09/2021

**Public title** 

Randomized, Open-Label, Phase II, Multicenter, Multi-Country Study to Evaluate Safety and Efficacy of Dasatinib 50 mg in First-Line Treatment of Early Chronic Phase Chronic Myeloid Leukemia

Scientific title

Randomized, Open-Label, Phase II, Multicenter, Multi-Country Study to Evaluate Safety and Efficacy of Dasatinib 50 mg in First-Line Treatment of Early Chronic Phase Chronic Myeloid Leukemia

Brief summary of the study: English

The primary endpoint to be measured during the study is the proportion of patients who achieve and maintain MMR at 12 months using RQ-PCR test. The study will be a multicenter, prospective, open-label, randomized Phase II study with a parallel design. Eligible patients with Ph+ CP CML will be randomly assigned to receive either dasatinib 50 mg QD or dasatinib 100 mg QD. The duration of patient participation will be 18 months.

Brief summary of the study: Arabic

Protocol number

LPI-JOR-LEB-KSA-TUN-2017-01

Study registered at the country of origin: Specify

Type of registration: Justify

N/A

Primary sponsor: Country of origin

Date of registration in national regulatory agency

05/11/2018

Acronym

NA

Acronym

NA



الهدف الأساسي من الدراسة هو قياس نسبة المرضى الذين يحققون استجابة جزي (MMR) جزينية كبرع (MMR) جزينية كبرع (MMR) جزينية كبرع (MMR) جزينية كبرع (MMR) بينية كبرع (MMR) المحلي (MMR) المحلي (MMR) المحلي المؤملين ال

Health conditions/problem studied: Specify

Early Chronic Phase Chronic Myeloid Leukemia

Interventions: Specify

dasatinib 50 mg QD or dasatinib 100 mg QD

Key inclusion and exclusion criteria: Inclusion criteria

Age ≥ 18 years.

Diagnosis of Ph+ or BCR-ABL positive CML in early CP (i.e. time from diagnosis <12 months).

Clonal evolution

ECOG performance of 0-2. Adequate end organ function

Key inclusion and exclusion criteria: Gender Key inclusion and exclusion criteria: Specify gender

Both

Key inclusion and exclusion criteria: Age minimum

Key inclusion and exclusion criteria: Age maximum

18 100

Key inclusion and exclusion criteria: Exclusion criteria

NYHA cardiac class 3-4 heart disease Cardiac symptoms

History of significant bleeding disorder

Patients with active uncontrolled psychiatric disorders

Pregnant or breast-feeding women

Patients in late chronic phase (i.e. time from diagnosis to treatment >12 months), accelerated phase (except as noted in inclusion criteria 2) or

blast phase

Type of study

Interventional

Type of intervention Type of intervention: Specify type

Pharmaceutical N/A

Trial scope Trial scope: Specify scope

Therapy

Study design: Allocation Study design: Masking

Randomized controlled trial Open (masking not used)

Study design: ControlStudy phaseDose comparison1

Study design: Purpose Study design: Specify purpose

Treatment

Study design: Assignment Study design: Specify assignment

Parallel N/A

IMP has market authorization IMP has market authorization: Specify



No

Name of IMP Year of authorization Month of authorization

Dasatinib

Type of IMP

Others

Pharmaceutical class

Tyrosine Kinase Inhibitor

Therapeutic indication

early chronic CML

Therapeutic benefit

Reduce the rate of adverse events and decrease cost of medications with the dose 50 mg while maintaining the efficacy.

enhance treatment compliance

Study model Study model: Explain model

N/A N/A

Study model: Specify model

N/A

Time perspective Time perspective: Explain time perspective

N/A

Time perspective: Specify perspective

N/A

Target follow-up duration Target follow-up duration: Unit

Number of groups/cohorts

Biospecimen retention Biospecimen description

NONE None retained

Target sample size Actual enrollment target size

100 100

Date of first enrollment: Type Date of first enrollment: Date

Actual 07/03/2019

Date of study closure: Type Date of study closure: Date

30/06/2022 Actual



Recruitment status Recruiting	Recruitment status: Specify	
Date of completion 30/06/2022		
IPD sharing statement plan	IPD sharing statement description	
No	NONE	
Additional data URL NA		
Admin comments		
Trial status		
Approved		
Secondary Identifying Numbers		
No Numbers		
Sources of Monetary or Material Suppor	t	
No Sources		
Secondary Sponsors		
No Sponsors		
Contact for Public/Scientific Queries		
No Contacts		



Centers/Hospitals Involved in the Study	
No Centers/Hospitals	
Ethics Review	
No Reviews	
Countries of Recruitment	
No Countries	
Health Conditions or Problems Studied	
No Problems Studied	
Interventions	
No Interventions	
Primary Outcomes	
No Outcomes	
Key Secondary Outcomes	
No Outcomes	



Trial Results	
Summary results	
Study results globally	
Date of posting of results summaries	Date of first journal publication of results
Results URL link	
Baseline characteristics	
Participant flow	
Adverse events	
Outcome measures	
URL to protocol files	