REPUBLIC OF LEBANON Lebanon Clinical Trials Registry

A Phase 3, Multi-center, Open-label, Randomized Study of Oral ABL001 Versus Bosutinib in Patients With Chronic Myelogenous Leukemia in Chronic Phase (CML-CP), Previously Treated With 2 or More Tyrosine Kinase Inhibitors

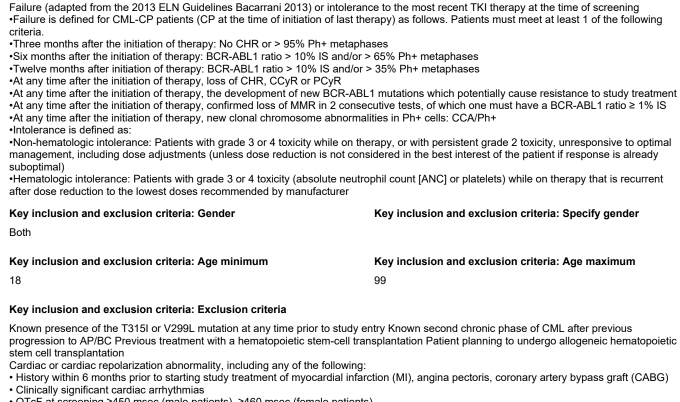
10/09/2025 18:17:51

| Primary registry identifying number | Protocol number |
|---|--|
| LBCTR2019010185 | CABL001A2301 |
| NOU registration number | |
| MOH registration number | |
| 49983/2017 | |
| Study registered at the country of origin | Study registered at the country of origin: Specify |
| Yes | |
| Type of registration | Type of registration: Justify |
| Retrospective | LCTR was already initiated, original file was previously submitted |
| Date of registration in national regulatory | |
| agency | |
| 21/12/2017 | |
| Primary sponsor | Primary sponsor: Country of origin |
| Novartis Pharma Services Inc. | Novartis Pharmaceuticals |
| Date of registration in primary registry | Date of registration in national regulatory agency |
| 14/08/2019 | 21/12/2017 |
| Public title | Acronym |
| A Phase 3, Multi-center, Open-label, Randomized Study of Oral | ASCEMBL |
| ABL001 Versus Bosutinib in Patients With Chronic Myelogenous Leukemia in Chronic Phase (CML-CP), Previously Treated With 2 | |
| or More Tyrosine Kinase Inhibitors | |
| Scientific title | Acronym |
| A Phase 3, Multi-center, Open-label, Randomized Study of Oral | |
| ABL001 Versus Bosutinib in Patients With Chronic Myelogenous Leukemia in Chronic Phase (CML-CP), Previously Treated With 2 | |
| or More Tyrosine Kinase Inhibitors | |
| Brief summary of the study: English | |
| The purpose of this pivotal study is to compare the efficacy of | |
| ABL001 with that of bosutinib in the treatment of patients with CML- CP having previously been treated with a minimum of two prior ATP | |
| -binding site TKIs with BCR-ABL ratios \geq 1% IS at screening. | |
| Brief summary of the study: Arabic | |
| ب لدى المرضى ABL001 مفتوحة اللصاقة، متعددة المراكز حول دواء3دراسة جزافيّة في المرحلة بن بسرطان الدم النقوي المزمن في المرحلة المزمنة، المعالجين سابعًا بمتبّطَيْن أو أكثر لكيناز التيروزيز | |
| Health conditions/problem studied: Specify | |
| Chronic Myelogenous Leukemia | |

ABL001. Bosutinib

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• QTcF at screening ≥450 msec (male patients), ≥460 msec (female patients)

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Male or female patients with a diagnosis of CML-CP ≥ 18 years of age

•< 30% blasts plus promyelocytes in peripheral blood and bone marrow</p>

Patients must meet all of the following laboratory values at the screening visit:

•Transient prior therapy related thrombocytopenia (< 50,000/mm3 for ≤ 30 days prior to screening) is acceptable

Prior treatment with a minimum of 2 prior ATP-binding site TKIs (i.e. imatinib, nilotinib, dasatinib, radotinib or ponatinib)

•No evidence of extramedullary leukemic involvement, with the exception of hepatosplenomegaly

BCR-ABL1 ratio ≥ 1% IS according to central laboratory at the screening examination

Key inclusion and exclusion criteria: Inclusion criteria

•< 15% blasts in peripheral blood and bone marrow</p>

< 20% basophils in the peripheral blood •≥ 50 x 109/L (≥ 50,000/mm3) platelets

· Long QT syndrome, family history of idiopathic sudden death or congenital long QT syndrome, or any of the following:

Risk factors for Torsades de Pointes (TdP)

· Concomitant medication(s) with a known risk of Torsades de Pointes per www.qtdrugs.org that cannot be discontinued or replaced 7 days prior to starting study drug by safe alternative medication.

- Inability to determine the QTcF interval
- · Severe and/or uncontrolled concurrent medical disease
- · History of acute pancreatitis within 1 year of study entry or past medical history of chronic pancreatitis
- · History of acute or chronic liver disease
- Treatment with medications that meet one of the following criteria and that cannot be discontinued at least one week prior to the start of
- treatment with study treatment
- Moderate or strong inducers of CYP3A Moderate or strong inhibitors of CYP3A and/or P-gp

• Women of child-bearing potential, unless they are using highly effective methods of contraception during dosing and for 3 days after last dose of ABL001.

 Sexually active males unless they use a condom during intercourse while taking the drug during treatment and for 3 days after stopping treatment and should not father a child in this period. A condom is required to be used also by vasectomized men as well as during intercourse with a male partner in order to prevent delivery of the drug via semen.

Type of study

Interventional

Type of intervention

Pharmaceutical

Type of intervention: Specify type

N/A

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| Trial scope Therapy | Trial scope: Specify scope N/A |
|--|---|
| Study design: Allocation Randomized controlled trial | Study design: Masking Open (masking not used) |
| Study design: Control Active | Study phase 3 |
| Study design: Purpose Treatment | Study design: Specify purpose N/A |
| Study design: Assignment Other | Study design: Specify assignment 2:1 |
| IMP has market authorization No | IMP has market authorization: Specify |
| Name of IMP ABL001 | Year of authorization Month of authorization |
| Type of IMP Cell therapy | |
| Pharmaceutical class orally bioavailable specific BCR-ABL inhibitor with a novel mechanism of ac | tion. |
| Therapeutic indication patients with Chronic Myelogenous Leukemia-CP who had prior treatment v binding site TKIs | vith two or more ATP |
| Therapeutic benefit increase OS & PFS | |
| Study model N/A | Study model: Explain model N/A |
| Study model: Specify model N/A | |
| Time perspective N/A Time perspective: Specify perspective N/A | Time perspective: Explain time perspective N/A |
| Target follow-up duration | Target follow-up duration: Unit |
| Number of groups/cohorts | |
| Biospecimen retention | Biospecimen description |



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| Samples without DNA | Bone marrow aspirate samples, Hematology , chemistry , coagulation, hepatitis , Liver function tests , are sent to Covance central laboratory, Navigate biopharma, molecular MD and Histogene X . |
|---|---|
| Target sample size | Actual enrollment target size |
| 5 | 3 |
| Date of first enrollment: Type Actual | Date of first enrollment: Date 05/09/2018 |
| Date of study closure: Type Actual | Date of study closure: Date 21/12/2022 |
| Recruitment status Recruiting | Recruitment status: Specify |
| Date of completion 29/02/2020 | |
| IPD sharing statement plan | IPD sharing statement description |
| Yes | Novartis is committed to sharing with qualified external researchers, access to patient-level data and supporting clinical documents from eligible studies. These requests are reviewed and approved by an independent review panel on the basis of scientific merit. All data provided is anonymized to respect the privacy of patients who have participated in the trial in line with applicable laws and regulations. |
| Additional data URL | |
| https://clinicaltrials.gov/ct2/show/record/NCT03106779?id=cabl001a2301&ra | ank=1 |

Admin comments

Trial status

Approved

| Secondary Identifying Numbers | | |
|--------------------------------|------------------------------|--|
| Full name of issuing authority | Secondary identifying number | |
| Clinical Trials.Gov | NCT03106779 | |

Sources of Monetary or Material Support

Name

Novartis Pharma Services Inc.



Secondary Sponsors

Name

NA

| Contact for Public/Scientific Queries | | | | | | |
|---------------------------------------|-------------------|------------|---------|------------------------------|-----------------------------------|--|
| Contact type | Contact full name | Address | Country | Telephone | Email | Affiliation |
| Public | Ali Bazarbachi | Beirut | Lebanon | 009613612 434 | bazarbac@aub.e du.lb | American University of Beirut Medical Center |
| Scientific | Hind Khairallah | Beirut | Lebanon | +961 1 512002 Ext. 271 | Hind.Khairallah@ fattal.com.lb | Khalil Fattal et Fils s.a.l. |
| Public | Joseph Kattan | Beirut | Lebanon | 009613635 913 | jkattan62@hotm ail.com | Hotel Dieu De France |
| Public | Dany ABi Gerges | Mansourieh | Lebanon | 009613341 960 | abgerges@idm.n et.lb | Bellevue Medical Center |

| Centers/Hospitals Involved in the Study | | | |
|--|---|---------------------|------------------|
| Center/Hospital name | me Name of principles investigator Principles investigator speciality Ethical approva | | Ethical approval |
| Bellevue Medical Center | Dr Dany Abi Gerges | Hematology Oncology | Approved |
| American University of Beirut Medical Center | Dr. Ali Bazarbachi | Hematology Oncology | Approved |
| Hotel Dieu De France | Dr Joseph Kattan | Hematology Oncology | Approved |

| Ethics Review | | | | |
|---|---------------|-----------------|-----------------------|--------------------------------|
| Ethics approval obtained | Approval date | Contact name | Contact email | Contact phone |
| American University of Beirut Medical Center | 05/06/2018 | Fuad Ziyadeh | fz05@aub.edu.lb | +961 (0) 1 350 000 ext:5445 |
| Hotel Dieu de France | 02/10/2017 | Nancy Alam | nancy.alam@usj.edu.lb | +961 1421000 ext 2335 |
| Bellevue Medical Center | 23/11/2017 | Ghassan Maalouf | gmaalouf@bmc.com.lb | +961 1 682666 ext 7600 |





| Countries of Recruitment |
|--------------------------|
| Name |
| Lebanon |
| Argentina |
| Australia |
| Belgium |
| Bulgaria |
| Canada |
| Czech Republic |
| France |
| Germany |
| Hungary |
| Italy |
| Japan |
| Republic of Korea |
| Netherlands |
| Turkey |
| United States of America |
| Saudi Arabia |
| United Kingdom |

| Health Conditions or Problems Studied | | |
|---------------------------------------|--------------------------------|-----|
| Condition Code Keyword | | |
| Chronic Myelogenous Leukemia | Leukaemia, unspecified (C95.9) | CML |





| Interventions | | | |
|---|---|--------------------|--|
| Intervention | Description | Keyword | |
| Physical examination, Vital Sign, Height and weight, ECOG performance status, Laboratory chemistry and hematology, Serology, Electrocardiogram (ECG), Echocardiogram, Pulmonary function tests, PK sampling (full/sparse), Bone Marrow Biopsy, Patient Report Outcomes (MDASI-CML, PGIC, WPAI, EQ5D-5L, resource | ICF, Lab tests, physical examination, ECG | Lab, ECG, ICF, BMA | |

| Primary Outcomes | | |
|-------------------------------------|-------------|---------|
| Name | Time Points | Measure |
| Major Molecular Response (MMR) rate | 24 weeks | 24 wks |

| Key Secondary Outcomes | | | |
|-------------------------------------|---|---------------------------|--|
| Name | Time Points | Measure | |
| Major Molecular Response (MMR) rate | 96 weeks after the last patient received the first study dose | 96 weeks after first dose | |
| Complete Cytogenetic response rate | 24,48,96 weeks | 24,48,96 weeks | |



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