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A Study to Investigate Tolerability and Efficacy of Asciminib (Oral) Versus Nilotinib (Oral) in Adult Participants (≥18 Years of Age) With Newly Diagnosed Philadelphia Chromosome Positive Chronic Myelogenous Leukemia in Chronic Phase (Ph+ CML-CP)

	12/08/2025 01:31:3
Main Information	
Primary registry identifying number	Protocol number
LBCTR2023025267	CABL001J12302
MOH registration number	
Study registered at the country of origin	Study registered at the country of origin: Specify
Yes	
Type of registration	Type of registration: Justify
Prospective	N/A
Date of registration in national regulatory agency	
Primary sponsor	Primary sponsor: Country of origin
Novartis Pharma AG	Novartis Pharma AG
Date of registration in primary registry	Date of registration in national regulatory agency
13/09/2023	
Public title	Acronym
A Study to Investigate Tolerability and Efficacy of Asciminib (Oral) Versus Nilotinib (Oral) in Adult Participants (≥18 Years of Age) With Newly Diagnosed Philadelphia Chromosome Positive Chronic Myelogenous Leukemia in Chronic Phase (Ph+ CML-CP)	
Scientific title	Acronym
A Phase IIIb, Multi-center, Open-label, Randomized Study of Tolerability and Efficacy of Oral Asciminib Versus Nilotinib in Patients With Newly Diagnosed Philadelphia Chromosome Positive Chronic Myelogenous Leukemia in Chronic Phase	ASC4START
Brief summary of the study: English	
The study is designed to compare the tolerability of asciminib versus nilotinib for the treatment of newly diagnosed, previously untreated patients with Positive Chronic Myelogenous Leukemia in Chronic Phase (Ph+ CML-CP)	
Brief summary of the study: Arabic	
مقابل نيلوتينيب لدى المرضى3دراسة متعدّدة المراكز ومفتوحة التسمية وعشوانيّة التوزيع في المرحلة تشخيص إصابتهم حديثًا بسرطان الدم النقوي المزمن الإيجابي لكروموسوم فيلادلفيا في المرحلة المزمنة	
Health conditions/problem studied: Specify	
Philadelphia Chromosome-Positive Chronic Myeloid Leukemia	

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Key inclusion and exclusion criteria: Specify gender

Key inclusion and exclusion criteria: Age maximum

Interventions: Specify

Drug: Asciminib Asciminib 80 mg QD administered under fasting conditions Other Name: ABL001 Drug: Nilotinib Nilotinib 300 mg BID administered under fasting conditions

Key inclusion and exclusion criteria: Inclusion criteria

Inclusion Criteria:

1- Patients with CML-CP within 3 months of diagnosis.

2- Diagnosis of CML-CP (ELN 2020 criteria) with cytogenetic confirmation of the Philadelphia chromosome

Documented chronic phase CML will meet all the below criteria Baccarani et al 2013:

< 15% blasts in peripheral blood and bone marrow,

< 30% blasts plus promyelocytes in peripheral blood and bone marrow,

< 20% basophils in the peripheral blood,

PLT count \geq 100 x 10^9/L (\geq 100,000/mm3), except treatment induced thrombocytopenia

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

3- Evidence of typical BCR::ABL1 transcript [e14a2 and/or e13a2] which is amenable to standardized RQ-PCR quantification by the central laboratory assessment.

4- ECOG performance status of 0 or 1.

5- Adequate end organ function as defined by:

Total bilirubin (TBL) < 3 x ULN; patients with Gilbert's syndrome may only be included if TBL \leq 3.0 x ULN or direct bilirubin \leq 1.5 x ULN, CrCl \geq 30 mL/min as calculated using Cockcroft-Gault formula, Serum lipase \leq 1.5 x ULN. For serum lipase > ULN - \leq 1.5 x ULN, value must be considered not clinically significant and not associated with risk factors for acute pancreatitis.

6- Patients must have the following laboratory values within normal limits or corrected to within normal limits with supplements prior to randomization:

Potassium (potassium increase of up to 6.0 mmol/L is acceptable if associated with CrCl* ≥ 90 mL/min),

Total calcium (corrected for serum albumin); (calcium increase of up to 12.5 mg/dl or 3.1 mmol/L is acceptable if associated with $CrCl^* \ge 90$ mL/min),

Magnesium (magnesium increase of up to 3.0 mg/dL or 1.23 mmol/L if associated with CrCl* \geq 90 mL/min), For patients with mild to moderate renal impairment (CrCl* \geq 30 mL/min and <90 mL/min) - potassium, total calcium (corrected for serum albumin) and magnesium should be within normal limits or corrected to within normal limits with supplements prior to randomization.

CrCl as calculated using Cockcroft-Gault formula. Other protocol-defined Inclusion/exclusion criteria will apply

Key inclusion and exclusion criteria: Gender

Both

Key inclusion and exclusion criteria: Age minimum

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Key inclusion and exclusion criteria: Exclusion criteria

Exclusion Criteria:

 Previous treatment of CML with any other anticancer agents including chemotherapy and/or biologic agents or prior stem cell transplant, with the exception of hydroxyurea and/or anagrelide.

 Known cytopathologically confirmed CNS infiltration (in absence of suspicion of CNS involvement, lumbar puncture not required).

3- Impaired cardiac function or cardiac repolarization abnormality including but not limited to any one of the following:

History of myocardial infarction (MI), angina pectoris, coronary artery bypass graft (CABG) within 6 months prior to starting study treatment. Clinically significant cardiac arrhythmias (e.g., ventricular tachycardia), complete left bundle branch block, high-grade AV block (e.g., bifascicular block, Mobitz type II and third degree AV block).

QTcF \ge 450 ms (male patients), \ge 460 ms (female patients) on the average of three serial baseline ECG (using the QTcF formula). If QTcF \ge 450 ms and electrolytes are not within normal ranges, electrolytes should be corrected and then the patient re-screened for QTcF. Long QT syndrome, family history of idiopathic sudden death or congenital long QT syndrome, or any of the following:

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Risk factors for Torsades de Pointes (TdP) including uncorrected hypokalemia or hypomagnesemia, history of cardiac failure, or history of clinically significant/symptomatic bradycardia.

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

Inability to determine the QTcF interval.

4- Severe and/or uncontrolled concurrent medical disease that in the opinion of the

Investigator could cause unacceptable safety risks or compromise compliance with the protocol (e.g. uncontrolled diabetes, active or uncontrolled infection; uncontrolled arterial or pulmonary hypertension, uncontrolled clinically significant hyperlipidemia).

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- 5- History of significant congenital or acquired bleeding disorder unrelated to cancer.
- 6- Major surgery within 4 weeks prior to study entry or patients who have not recovered from prior surgery.

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- 7- History of other active malignancy within 3 years prior to study entry with the exception of previous or concomitant basal cell skin cancer and previous carcinoma in situ treated curatively.
- 8- History of acute pancreatitis within 1 year prior to randomization or medical history of chronic pancreatitis.
- History of chronic liver disease leading to severe hepatic impairment, or ongoing acute liver disease.
- 10- Known history of chronic Hepatitis B (HBV), or chronic Hepatitis C (HCV) infection. Testing for Hepatitis B surface antigen (HBs Ag) and Hepatitis B core antibody (HBc Ab/anti HBc) will be performed at screening. If anti-HBc is positive, HBV-DNA evaluation will be carried out at screening. A patient having positive HBV-DNA will not be enrolled in the study. Also, a patient with positive HBSAg will not be enrolled in the study. HCV Ab testing will also be performed at screening. For details on the criteria see Appendix 4.
- 11- History of Human Immunodeficiency Virus (HIV) unless well-controlled on a stable d dose of anti-retroviral therapy at the time of screening.

Other protocol-defined Inclusion/exclusion criteria will apply

Type of study

Interventional

Type of intervention	Type of intervention: Specify type
Pharmaceutical	N/A
Trial scope	Trial scope: Specify scope
Therapy	N/A
Study design: Allocation	Study design: Masking
Randomized controlled trial	Open (masking not used)
Study design: Control	Study phase
Active	3
Study design: Purpose	Study design: Specify purpose
Treatment	N/A
Study design: Assignment	Study design: Specify assignment
Parallel	N/A
IMP has market authorization No	IMP has market authorization: Specify
Name of IMP Asciminib	Year of authorization Month of authorization
Type of IMP Immunological	
Pharmaceutical class tyrosine kinase inhibitor (TKI)	

Therapeutic indication Philadelphia Chromosome-Positive Chronic Myeloid Leukemia

Therapeutic benefit Treatment

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Study model N/A	Study model: Explain model N/A
N/A	N/A
Study model: Specify model N/A	
Time perspective N/A	Time perspective: Explain time perspective
Time perspective: Specify perspective N/A	
Target follow-up duration	Target follow-up duration: Unit
Number of groups/cohorts	
Biospecimen retention	Biospecimen description
Samples with DNA**	Samples will be shipped to ICON Specialty Lab
Target sample size 5	Actual enrollment target size
Date of first enrollment: Type	Date of first enrollment: Date
Anticipated	17/04/2023
Date of study closure: Type	Date of study closure: Date
Anticipated	15/03/2027
Recruitment status Pending	Recruitment status: Specify
Date of completion	
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	This trial data availability is according to the criteria and process described on www.clinicalstudydatarequest.com

https://clinicaltrials.gov/ct2/show/record/NCT05456191?term=CABL001J12302&draw=2&rank=1





Admin comments

Trial status

Approved

Secondary Identifying Numbers		
Full name of issuing authority	Secondary identifying number	
Clinicaltrials.gov	NCT05147220	

Sources of Monetary or Material Support	
Name	
Novartis Pharma AG	

Secondary Sponsors
Name
NA

Contac	Contact for Public/Scientific Queries					
Contact type	Contact full name	Address	Country	Telephone	Email	Affiliation
Public	Ali Bararbachli	Beirut	Lebanon	+961 3 612434	bazarbac@aub.e du.lb	American University of Beirut Medical Center
Scientific	Hind Khairallah	Sin El Fil	Lebanon	009611512 002 Ext. 271 E	hind.khairallah@f attal.com.lb	Khalil Fattal et Fils s.a.l

Centers/Hospitals Involved in the Study				
Center/Hospital name	Name of principles investigator	Principles investigator speciality	Ethical approval	
American University of Beirut Medical Center	Ali Bazarbachli	Oncology - Hematology	Approved	



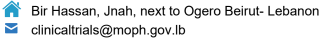


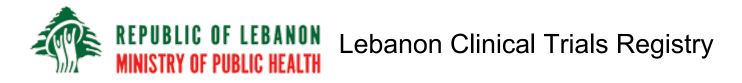
Ethics Review				
Ethics approval obtained	Approval date	Contact name	Contact email	Contact phone
American University of Beirut Medical Center	23/11/2022	Rami Mahfouz	rm11@aub.edu.lb	961 (0) 1 350 000 ext:5445

Countries of Recruitment
Name
France
Czech Republic
Germany
Hungary
Bulgaria
Slovakia
Canada
Greece
Italy
Malaysia
Netherlands
Oman
South Africa
Switzerland
United States of America

Health Conditions or Problems Studied

Condition	Code	Keyword
Philadelphia Chromosome-Positive Chronic Myeloid Leukemia	Chronic myeloid leukaemia (C92.1)	CML





Interventions			
Intervention	Description	Keyword	
Consenting, IMP administration, Laboratory testing, Imaging	Consenting, IMP administration, Laboratory testing, Imaging	Consenting, IMP administration, Laboratory testing, Imaging	

Primary Outcomes				
Name	Time Points	Measure		
Time to discontinuation of study treatment due to adverse event (TTDAE)	From date of first dose to date of treatment discontinuation due to AE, assessed up to 4.5 years	TTDAE is defined as the time from the date of first dose of study treatment to the date of discontinuation of study treatment due to adverse event (AE)		

Key Secondary Outcomes			
Name	Time Points	Measure	
Percentage of participants with Major Molecular response (MMR) at scheduled data collection time points	Screening, Week 4, Week 12 and thereafter every 12 weeks until End of Treatment (EOT) and EOT, assessed up to approximately 4.5 years	MMR will be assessed using fusion gene of the BCR and ABL genes (BCR-ABL) transcript levels measured by realtime quantitative polymerase chain reaction. The percentage of participants with MMR at each time point will be assessed	
Percentage of participants with Major Molecular response (MMR) by scheduled data collection time points	Screening, Week 4, Week 12 and thereafter every 12 weeks until End of Treatment (EOT) and EOT, assessed up to approximately 4.5 years	MMR will be assessed using BCR-ABL transcript levels measured by realtime quantitative polymerase chain reaction. The percentage of participants who meet the criteria for having achieved the endpoint (MMR) at or before the specified visit will be calculated	
Percentage of participants with MR4.0 at scheduled data collection time points	Screening, Week 4, Week 12 and thereafter every 12 weeks until End of Treatment (EOT) and EOT, assessed up to approximately 4.5 years	MR4.0 will be assessed using BCR-ABL transcript levels measured by realtime quantitative polymerase chain reaction. The percentage of participants with MR4.0 at each time point will be assessed	
Percentage of participants with MR4.0 by scheduled data collection time points	Screening, week 4, week 12 and thereafter every 12 weeks until End of Treatment (EOT) and EOT, assessed up to approximately 4.5 years	MR4.0 will be assessed using BCR-ABL transcript levels measured by realtime quantitative polymerase chain reaction. The percentage of participants who meet the criteria for having achieved the endpoint (MR4.0) at or before the specified visit will be calculated	
Percentage of participants with MR4.5 at scheduled data collection time points	Screening, Week 4, Week 12 and thereafter every 12 weeks until End of Treatment (EOT) and EOT, assessed up to approximately 4.5 years	MR4.5 will be assessed using BCR-ABL transcript levels measured by realtime quantitative polymerase chain reaction. The percentage of participants with MR4.5 at each time point will be assessed	
Percentage of participants with MR4.5 by scheduled data collection time points	Screening, Week 4, Week 12 and thereafter every 12 weeks until End of Treatment (EOT) and EOT, assessed up to approximately 4.5 years	MR4.5 will be assessed using BCR-ABL transcript levels measured by realtime quantitative polymerase chain reaction. The percentage of participants who meet the criteria for having achieved the endpoint (MR4.5) at or before the specified visit will be calculated	
Percentage of participants with Complete Hematological response (CHR) at scheduled data collection time points	Screening, Week 4, Week 12 and thereafter every 12 weeks until End of Treatment (EOT) and EOT, assessed up to approximately 4.5 years	Hematologic response will be assessed by complete blood count and physical examination at each visit. The percentage of participants with CHR at each time point will be assessed	

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Percentage of participants with Complete Hematological response (CHR) by scheduled data collection time points	Screening, Week 4, Week 12 and thereafter every 12 weeks until End of Treatment (EOT) and EOT, assessed up to approximately 4.5 years	Hematologic response will be assessed by complete blood count and physical examination at each visit. The percentage of participants who meet the criteria for having achieved the endpoint (CHR) at or before the specified visit will be calculated
Percentage of participants with BCR::ABL1 ratio ≤1% by Week 48 and Week 96	Week 48 and Week 96	The percentage of participants who meet the criteria for having achieved BCR::ABL1 ratio ≤1% at or before the specified visit will be calculated
Duration of MMR	From the date of the first documented molecular response at MMR level to the date of first documented loss of MMR or death due to any cause, whichever occurs first, assessed up to approximately 4.5 years	Duration of MMR is defined as the time between the date of the first documented achievement MMR and the earliest date of loss of MMR, treatment failure, progression to AP/BC, or CML-related death
Duration of MR4.0	From the date of the first documented molecular response at MR4 level to the date of first documented loss of the response level or death due to any cause, whichever occurs first, assessed up to approximately 4.5 years	Duration of MR4.0 is defined as the time between the date of the first documented achievement MR4 and the earliest date of loss of MR4, treatment failure, progression to AP/BC, or CML-related death
Duration of MR4.5	From the date of the first documented molecular response at MR4.5 level to the date of first documented loss of the response level or death due to any cause, whichever occurs first, assessed up to approximately 4.5 years	Duration of MR4.5 is defined as the time between the date of the first documented achievement MR4.5 and the earliest date of loss of MR4.5, treatment failure, progression to AP/BC, or CML-related death
Time to first MMR	From the date of randomization to the date of the first MMR, assessed up to approximately 4.5 years	Time to first MMR is defined as the time from the date of randomization to the date of the first documented occurrence of MMR
Time to first MR4.0	From the date of randomization to the date of the first MR4, assessed up to approximately 4.5 years	Time to first MR4.0 is defined as the time from the date of randomization to the date of the first documented occurrence of MR4
Time to first MR4.5	From the date of randomization to the date of the first MR4.5, assessed up to approximately 4.5 years	Time to first MR4.5 is defined as the time from the date of randomization to the date of the first documented occurrence of MR4.5
Time to treatment failure (TTF)	Up to approximately 4.5 years	TTF is defined as the time from date of randomization to the first/earliest documented date of any of the following events: Treatment failure per European leukemia network (ELN) criteria, Confirmed loss of MMR (in 2 consecutive tests) at any time while on study treatment, Discontinuation from study treatment due to any reason
Event free survival (EFS)	Up to approximately 4.5 years	EFS is defined as the time from the date of the first dose of study treatment to the earliest occurrence of treatment failure, confirmed lost of MMR, discontinuation due to AE, progression to AP/BC, and death from any cause
Progression free survival (PFS)	Up to approximately 4.5 years	PFS is defined as the time from the date of randomization to the earliest occurrence of progression to AP/BC or death from any cause
Overall survival (OS)	Up to approximately 4.5 years	OS is defined as the time from the date of randomization to the date of death from any cause
Time to treatment discontinuation (TTD) due to selected reasons	Up to approximately 4.5 years	TTD is the time from the date of first dose of study treatment to the date of discontinuation of study treatment due to lack of efficacy, treatment failure, disease progression, suboptimal response or death



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Change from baseline in overall scores and individual scales of the European organization for research and treatment of cancer - quality of life questionnaire (EORTC QLQ-C30)	Baseline, every 4 weeks from Week 4 to Week 12, after Week 24, Week 48, Week 96, EOT and every 4 weeks until 12 weeks after EOT, assessed up to approximately 4.5 years	Change from baseline in Overall Scores and individual domains of the EORTC QLQ-C30. The EORTC QLQ- C30 contains 30 items and is composed of both multi- item scales and single-item measures based on the participant's experience over the past week. These include five functional scales (physical, role, emotional, cognitive and social functioning), three symptom scales (fatigue, nausea/vomiting, and pain), six single items (dyspnea, insomnia, appetite loss, constipation, diarrhea and financial impact) and a global health status/QoL scale
Change from baseline in overall scores and individual scales of the European organization for research and treatment of cancer CML module (EORTC QLQ-CML24)	Baseline, every 4 weeks from Week 4 to Week 12, after Week 24, Week 48, Week 96, EOT and every 4 weeks until 12 weeks after EOT, assessed up to approximately 4,5 years	Change from baseline in Overall Scores and individual domains of the EORTC QLQ-CML24. The EORTC QLQ-CML24 assesses specific concepts relevant to the experience of patients with CML. The QLQ-CML24 has 24 items which assess symptom burden, impact on daily life and on worry/mood, body image problems, and satisfaction with care and with social life based on the participant's experience over the past week

Trial Results

Summary results

Study results globally

Date of posting of results summaries

Results URL link

Baseline characteristics

Participant flow

Adverse events

Outcome measures

URL to protocol files

Date of first journal publication of results