



Roll-over Study to Allow Continued Access to Ribociclib

12/08/2025 18:20:13

Main Information

Primary registry identifying number

LBCTR2022095114

Protocol number

CLEE011A2412B

MOH registration number

Study registered at the country of origin

Yes

Study registered at the country of origin: Specify

Type of registration

Prospective

Type of registration: Justify

N/A

Date of registration in national regulatory agency

Primary sponsor

Novartis Pharmaceuticals

Primary sponsor: Country of origin

Novartis Pharmaceuticals

Date of registration in primary registry

01/08/2023

Date of registration in national regulatory agency

Public title

Roll-over Study to Allow Continued Access to Ribociclib

Acronym

Scientific title

A Post-trial Access Roll-over Study to Allow Access to Ribociclib (LEE011) for Patients Who Are on Ribociclib Treatment in Novartis-sponsored Study

Acronym

Brief summary of the study: English

This is an open-label, multi-center, roll-over study to evaluate the long term safety of ribociclib in combination with other drugs in participants who are participating in a Novartis sponsored global study, that has fulfilled requirements for its primary objective(s), and who in the opinion of the Investigator, would benefit from continued treatment.

Brief summary of the study: Arabic

هذه دراسة تمديد مفتوحة التسمية ومتعددة المراكز لتقييم السلامة طويلة المدى لريبوسيكليب بالاشتراك مع أدوية أخرى في المشاركين الذين يشاركون في دراسة عالمية ترعاها شركة نوفارتيس، والتي أوفت بمتطلبات هدفها الأساسي (أهدافها)، ومن يعتقد طبيب الدراسة أنه سيستفيد من استمرار العلاج.

Health conditions/problem studied: Specify

Metastatic Breast Cancer

Interventions: Specify

Drug: Ribociclib (Participants continue ribociclib as was administered in their parent study)
Drug: Letrozole (Participants continue ribociclib in combination with letrozole as was administered in their parent study)
Drug: Anastrozole (Participants continue ribociclib in combination with anastrozole as was administered in their parent study)
Drug: Goserelin (Participants continue ribociclib in combination with goserelin as was administered in their parent study)
Drug: Tamoxifen (Participants continue ribociclib in combination with tamoxifen as was administered in their parent study)
Drug: Fulvestrant (All participants continue ribociclib in combination with fulvestrant as was administered in their parent study)

Key inclusion and exclusion criteria: Inclusion criteria





1. Currently participating in a Novartis sponsored global study (parent study), receiving treatment with ribociclib in combination with other drugs, and the parent study has fulfilled its primary objective(s)
2. Must have been receiving treatment with ribociclib for at least 6 cycles in the parent study
3. Currently has evidence of clinical benefit as determined by the Investigator

Key inclusion and exclusion criteria: Gender

Both

Key inclusion and exclusion criteria: Specify gender**Key inclusion and exclusion criteria: Age minimum**

18

Key inclusion and exclusion criteria: Age maximum

99

Key inclusion and exclusion criteria: Exclusion criteria

1. Permanent discontinuation of ribociclib in the parent study
2. Currently has unresolved toxicities for which ribociclib dosing has been interrupted in the parent study
3. Local access to commercially available ribociclib and reimbursed

Type of study

Interventional

Type of intervention

Pharmaceutical

Type of intervention: Specify type

N/A

Trial scope

Therapy

Trial scope: Specify scope

N/A

Study design: Allocation

N/A

Study design: Masking

Open (masking not used)

Study design: Control

Uncontrolled

Study phase

4

Study design: Purpose

Treatment

Study design: Specify purpose

N/A

Study design: Assignment

Single

Study design: Specify assignment

N/A

IMP has market authorization

Yes, Lebanon and Worldwide

IMP has market authorization: Specify

US, EU, and other countries

Name of IMP

Ribociclib

Year of authorization

2017

Month of authorization

1

Type of IMP

Others

Pharmaceutical class

highly selective small molecule inhibitor of cyclin-dependent kinases 4 and 6 (CDK4/6)

Therapeutic indication

Metastatic Breast Cancer

Therapeutic benefit

continued treatment to participants who are currently receiving ribociclib

Study model

N/A

Study model: Explain model

Study model: Specify model

N/A

N/A

Time perspective

Time perspective: Explain time perspective

N/A

N/A

Time perspective: Specify perspective

N/A

Target follow-up duration

Target follow-up duration: Unit

Number of groups/cohorts

Biospecimen retention

None retained

Biospecimen description

NA

Target sample size

4

Actual enrollment target size

Date of first enrollment: Type

Anticipated

Date of first enrollment: Date

31/10/2022

Date of study closure: Type

Anticipated

Date of study closure: Date

16/02/2028

Recruitment status

Pending

Recruitment status: Specify

Date of completion

IPD sharing statement plan

Yes

IPD sharing statement description

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.

This trial data availability is according to the criteria and process described on www.clinicalstudydatarequest.com

Additional data URL

<https://clinicaltrials.gov/ct2/show/record/NCT05161195?term=clee011A2412B&draw=2&rank=1>

Admin comments

**Trial status**

Approved

Secondary Identifying Numbers

| Full name of issuing authority | Secondary identifying number |
|--------------------------------|------------------------------|
| clinicaltrials.gov | NCT05161195 |

Sources of Monetary or Material Support

| Name |
|--------------------------|
| Novartis Pharmaceuticals |

Secondary Sponsors

| Name |
|------|
| NA |

Contact for Public/Scientific Queries

| Contact type | Contact full name | Address | Country | Telephone | Email | Affiliation |
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| Public | Joseph Kattan | Beirut | Lebanon | +961 3 635913 | jkattan62@hotmail.com | Hotel Dieu de France |



Centers/Hospitals Involved in the Study

| Center/Hospital name | Name of principles investigator | Principles investigator speciality | Ethical approval |
|--|---------------------------------|------------------------------------|------------------|
| American University of Beirut Medical Center | Nagi El Saghir | Hematology Oncology | Approved |
| Ain Wazein Medical Village | Jawad Makarem | Hematology Oncology | Approved |
| Hotel Dieu de France | Joseph Kattan | Hematology Oncology | Approved |

Ethics Review

| Ethics approval obtained | Approval date | Contact name | Contact email | Contact phone |
|--|---------------|------------------|--------------------------|-----------------------------|
| American University of Beirut Medical Center | 05/09/2022 | Rami Mahfouz | rm11@aub.edu.lb | +961 (0) 1 350 000 ext:5445 |
| Ain w Zein Medical Village | 25/08/2022 | Hayat Kamaledine | irb@awmedicalvillage.org | +961 (0) 5 509 001 ext 2014 |
| Hotel Dieu de France | 12/08/2022 | Nancy Alam | nancy.alam@usj.edu.lb | +961 (0) 1 421000 ext 2335 |



Countries of Recruitment

| Name |
|--------------------------|
| Lebanon |
| Poland |
| Singapore |
| United States of America |
| Brazil |
| France |
| Greece |
| China |
| Italy |
| Japan |
| Republic of Korea |
| Mexico |
| Portugal |
| South Africa |
| Spain |
| Taiwan |
| Turkey |

Health Conditions or Problems Studied

| Condition | Code | Keyword |
|--------------------------|------------------------------------|--------------------------|
| Metastatic Breast Cancer | Malignant neoplasm of breast (C50) | Metastatic Breast Cancer |

Interventions

| Intervention | Description | Keyword |
|-------------------------------------|-------------------------------------|-------------------------------------|
| ICF, IMP administration, local Labs | ICF, IMP administration, local Labs | ICF, IMP administration, local Labs |



Primary Outcomes

| Name | Time Points | Measure |
|---|--|--|
| Percentage of participants with treatment-emergent adverse events (AES) | From day of first dose of study medication to 30 days after last dose of study medication, up to 5 years | The percentage of participants with treatment-emergent adverse events will be summarized, including significant adverse events leading to discontinuation, and adverse events leading to dose adjustment |

Key Secondary Outcomes

| Name | Time Points | Measure |
|-----------------------|---------------|--|
| Clinical benefit rate | Up to 5 years | Percentage of participants with clinical benefit as assessed by the Investigator at scheduled study visits |

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