## REPUBLIC OF LEBANON Lebanon Clinical Trials Registry

COMPLEEMENT-1: An Open-label, Multicenter, Phase IIIb Study to Assess the Safety and Efficacy of Ribociclib (LEE011) in Combination With Letrozole for the Treatment of Men and Pre/Postmenopausal Women With Hormone Receptor-positive (HR+) HER2-negative (HER2-) Advanced Breast Cancer (aBC) With no Prior Hormonal Therapy for Advanced Disease

14/08/2025 23:13:31 Main Information Primary registry identifying number **Protocol number** LBCTR2019010184 CLEE011A2404 MOH registration number 20521/2017 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 by Paper Date of registration in national regulatory agency 01/06/2017 **Primary sponsor** Primary sponsor: Country of origin Novartis Pharma Services Inc. Novartis Pharmaceuticals Date of registration in national regulatory agency Date of registration in primary registry 27/01/2021 01/06/2017 Public title Acronym COMPLEEMENT-1: An Open-label, Multicenter, Phase IIIb Study to **COMPLEEMENT 1** Assess the Safety and Efficacy of Ribociclib (LEE011) in Combination With Letrozole for the Treatment of Men and Pre/Postmenopausal Women With Hormone Receptor-positive (HR+) HER2-negative (HER2-) Advanced Breast Cancer (aBC) With no Prior Hormonal Therapy for Advanced Disease Scientific title Acronym COMPLEEMENT-1: An Open-label, Multicenter, Phase IIIb Study to Assess the Safety and Efficacy of Ribociclib (LEE011) in Combination With Letrozole for the Treatment of Men and Pre/Postmenopausal Women With Hormone Receptor-positive (HR+) HER2-negative (HER2-) Advanced Breast Cancer (aBC) With no Prior Hormonal Therapy for Advanced Disease Brief summary of the study: English The purpose of this Phase IIIb study is to collect additional safety and efficacy data for the combination of ribociclib + letrozole in men and pre/postmenopausal women with HR+HER2- advanced breast cancer.

Brief summary of the study: Arabic

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Key inclusion and exclusion criteria: Specify gender

Key inclusion and exclusion criteria: Age maximum

بالاشتراك مع ليتروزول لعلاج (LEE011) دراسة مفتوحة اللصاقة، متعددة المراكز في المرحلة الثالثة ب لتقييم سلامة وفعاليّة رييوسيكليب الذين (-HR) 2وسلبيّ الهير (+HR) الرجال والنساء قبل/بعد انقطاع الطمث المصابين بسرطان الثدي المتقدّم الإيجابيّ مستقبلات الهرمون لم يتلقوا أي علاج هرمونيّ سابق للمرض المتقدّم

### Health conditions/problem studied: Specify

Advanced Breast Cancer

### Interventions: Specify

•Drug: Ribociclib •Drug: Letrozole •Drug: Goserelin

#### Key inclusion and exclusion criteria: Inclusion criteria

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MINISTRY OF PUBLIC HEALTH

•Male or female advanced (locoregionally recurrent or metastatic) breast cancer not amenable to curative therapy.

•In the case of women, both pre/perimenopausal and postmenopausal patients are eligible

•Patient has a histologically and/or cytologically confirmed diagnosis of estrogen-receptor positive and/or progesterone receptor positive breast cancer

•Patient has HER2-negative breast cancer defined as a negative in situ hybridization test or an IHC status of 0, 1+ or 2+. If IHC is 2+, a negative in situ hybridization (FISH, CISH, or SISH) test is required

•Patient has an Eastern Cooperative Oncology Group (ECOG) performance status ≤2

•Patient has adequate bone marrow and organ function as defined by ALL of the following laboratory values (as assessed by local laboratory): ∘Absolute neutrophil count ≥ 1.5 × 10^9/L

∘Platelets ≥ 100 × 10^9/L

∘Hemoglobin ≥ 9.0 g/dL

•Potassium, sodium, calcium corrected for serum albumin and magnesium within normal limits or corrected to within normal limits with supplements before first dose of the study medication

∘INR ≤1.5

∘Serum creatinine <1.5 mg/dl or creatinine clearance≥50 mL/min

In absence of liver metastases, alanine aminotransferase (ALT) and aspartate aminotransferase (AST) should be below 2.5 × ULN. If the patient has liver metastases, ALT and AST should be < 5 × ULN.</p>

• Total serum bilirubin < ULN; or total bilirubin ≤ 3.0 × ULN with direct bilirubin within normal range in patients with well-documented Gilbert's Syndrome

Patient must have a 12-lead ECG with ALL of the following parameters at screening:
 QTcF interval at screening <450 msec (using Fridericia's correction)</li>
 Resting heart rate ≥ 50 bpm

Key inclusion and exclusion criteria: Gender

Both

Key inclusion and exclusion criteria: Age minimum

18

### Key inclusion and exclusion criteria: Exclusion criteria

Patient who received any CDK4/6 inhibitor
Patient who received any prior systemic hormonal therapy for advanced breast cancer; no more than one prior regimen of chemotherapy for the treatment of metastatic disease is permitted

99

Type of study

Interventional

| <b>Type of intervention</b> | <b>Type of intervention: Specify type</b> |
|-----------------------------|---|
| Pharmaceutical              | N/A                                       |
| <b>Trial scope</b>          | Trial scope: Specify scope                |
| Therapy                     | N/A                                       |
| Study design: Allocation    | Study design: Masking                     |
| N/A: Single arm study       | Open (masking not used)                   |
| Study design: Control       | Study phase                               |

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| N/A  | 3   |   |
|--|---|---|
| Study design: Purpose  | Study design: Specify purpos  | se  |
| Treatment  | N/A   |   |
| Study design: Assignment   | Study design: Specify assigr  | nment   |
| Single   | N/A   |   |
| IMP has market authorization   | IMP has market authorization  | 1: Specify  |
| Yes, Lebanon and Worldwide   | USA, EU & other countries   | i. Opecity  |
|  |   |   |
| Name of IMP  | Year of authorization   | Month of authorization  |
| Ribociclib ( Kisqali)  | 2017  | 8   |
| Type of IMP  |   |   |
| Others   |   |   |
| Pharmaceutical class   |   |   |
| Orally bioavailable, highly selective small molecule inhibitor of cyclin-deperture (CDK4/6).   | endent kinases 4 and 6  |   |
| Therapeutic indication   |   |   |
| Men and Pre/Postmenopausal Women With Hormone Receptor-positive (<br>-) Advanced Breast Cancer | HR+) HER2-negative (HER2  |   |
| Therapeutic benefit  |   |   |
| increase Overall survival and progression free survival  |   |   |
| Study model  | Study model: Explain model  |   |
| N/A  | N/A   |   |
|  |   |   |
| Study model: Specify model   |   |   |
| N/A  |   |   |
|  |   |   |
| Time perspective   | Time perspective: Explain tin   | ne perspective  |
| N/A  | N/A   |   |
| Time perspective: Specify perspective  |   |   |
| N/A  |   |   |
|  |   |   |
|  |   |   |
| Target follow-up duration  | Target follow-up duration: Ur   | nit   |
|  |   |   |
| Number of groups/cohorts   |   |   |
|  |   |   |
|  | <b>D</b> iagonalizza in 1911  |   |
| Biospecimen retention  | Biospecimen description   | anlee will be obing ad autoide                                      |
| None retained  | Local lab is being used, no san<br>Lebanon, Lab samples are mai<br>patient safety | nples will be shipped outside<br>inly CBC, Chemistry to follow up c |
|  |   |   |
|  |   |   |

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| <b>Target sample size</b><br>20   | Actual enrollment target size<br>17   |
|---|---|
| Date of first enrollment: Type<br>Actual  | Date of first enrollment: Date  |
| Date of study closure: Type<br>Actual   | Date of study closure: Date<br>30/08/2021   |
| Recruitment status<br>Complete  | Recruitment status: Specify   |
| Date of completion<br>12/01/2018  |   |
| IPD sharing statement plan  | IPD sharing statement description   |
| Yes   | Novartis is committed to sharing with qualified external<br>researchers, access to patient-level data and supporting clinical<br>documents from eligible studies. These requests are reviewed<br>and approved by an independent review panel on the basis of<br>scientific merit. All data provided is anonymized to respect the<br>privacy of patients who have participated in the trial in line with<br>applicable laws and regulations. |
| Additional data URL   |   |
| https://clinicaltrials.gov/ct2/show/record/NCT02941926?<br>recrs=d&rslt=Without&type=Intr&cond=Advanced+Breast+Cancer&titles=cc | mpleement&spons=novartis&phase=2&rank=1   |
| Admin comments  |   |

**Trial status** 

Approved

| Secondary Identifying Numbers  |                              |
|--------------------------------|------------------------------|
| Full name of issuing authority | Secondary identifying number |
| Clinical Trials.Gov            | NCT02941926                  |

## Sources of Monetary or Material Support

### Name

Novartis Pharma Services Inc.

| Secondary Sponsors |  |
|--------------------|--|
| Name               |  |
| NA                 |  |



| Contact for Public/Scientific Queries |                   |            |         |                              |                                   |                                    |
|---------------------------------------|-------------------|------------|---------|------------------------------|-----------------------------------|------------------------------------|
| Contact<br>type                       | Contact full name | Address    | Country | Telephone                    | Email                             | Affiliation                        |
| Public                                | Fadi Farhat       | Saida      | Lebanon | +961 3<br>753 155            | drfadi.trials@gm<br>ail.com       | Hammoud<br>Hospital                |
| Scientific                            | Hind Khairallah   | Beirut     | Lebanon | +961 1<br>512002<br>Ext. 271 | Hind.Khairallah@<br>fattal.com.lb | Khalil<br>Fattal et<br>Fils s.a.l. |
| Public                                | Fadi El Karak     | Mansourieh | Lebanon | +961 3<br>061 621            | felkarak@yahoo.<br>com            | Bellevue<br>Medical<br>Center      |
| Public                                | Georges Chahine   | Beirut     | Lebanon | +9613<br>647778              | Chahine_george<br>s@hotmail.com   | Hotel Dieu<br>De France            |

| Centers/Hospitals Involved in the Study    |                                 |                                    |                  |  |
|--|---------------------------------|------------------------------------|------------------|--|
| Center/Hospital name                       | Name of principles investigator | Principles investigator speciality | Ethical approval |  |
| Hammoud Hospital University Medical Center | Dr Fadi Farhat                  | Hematology Oncology                | Approved         |  |
| Hotel Dieu De France                       | Dr Georges Chahine              | Hematology Oncology                | Approved         |  |
| Bellevue Medical Center                    | Dr Fadi El Karak                | Hematology Oncology                | Approved         |  |

| Ethics Review                                    |               |                 |                             |                              |
|--|---------------|-----------------|-----------------------------|------------------------------|
| Ethics approval obtained                         | Approval date | Contact name    | Contact email               | Contact phone                |
| Hotel Dieu de France                             | 02/05/2017    | Sami Richa      | cue@usj.edu.lb              | 961421229                    |
| Bellevue Medical<br>Center                       | 21/08/2017    | Ghassan Maalouf | Gmaalouf@bmc.com.lb         | 961 (0) 1 682666 ext<br>5006 |
| Hammoud Hospital<br>University Medical<br>Center | 02/05/2017    | Ahmad Zaatari   | zaatari@hammoudhospital.com | 961 (0) 7 723111 ext<br>1160 |





### **Countries of Recruitment**

| Name                     |
|--------------------------|
| Lebanon                  |
| Argentina                |
| Austria                  |
| Belgium                  |
| Bulgaria                 |
| Canada                   |
| Jordan                   |
| Oman                     |
| Saudi Arabia             |
| Spain                    |
| United Kingdom           |
| United States of America |

| Health Conditions or Problems Studied |                             |               |
|---------------------------------------|-----------------------------|---------------|
| Condition Code Keyword                |                             |               |
| Advanced Breast Cancer                | Breast, unspecified (C50.9) | Breast Cancer |

| Interventions   |  |                             |  |
|---|--|-----------------------------|--|
| Intervention  | Description                            | Keyword                     |  |
| Hematology tests: WBC, ANC, lymphocyte,<br>hemoglobin, platelets (as clinically indicated),<br>Chemistry tests: Alkaline phosphatase, ALT<br>(SGPT), AST (SGOT), calcium corrected for<br>serum albumin, creatinine or creatinine<br>clearance, potassium, sodium,<br>magnesium,direct bilirubin, total bilirubin (as<br>clinically indicated), ECG | Lab tests , ECG , Radiology assessment | Lab tests , Radiology , ECG |  |

| Primary Outcomes   |             |                           |  |
|--|-------------|---------------------------|--|
| Name   | Time Points | Measure                   |  |
| The number of participants with adverse events as a measure of safety and tolerability | PFS         | Progression free survival |  |



| Key Secondary Outcomes   |             |                           |
|--|-------------|---------------------------|
| Name   | Time Points | Measure                   |
| Time-to-Progression (TTP), Overall response rate (ORR), ,<br>Clinical Benefit Rate (CBR) | PFS         | Progression free survival |

| 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                |  |
|                                      |  |
|                                      |  |