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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

20/08/2025 14:54:12 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 17/12/2019 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)
 Resting heart rate ≥ 50 bpm

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

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

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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
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 purpo	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 (-) 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		
	D iagonalizza in 1911	
Biospecimen retention	Biospecimen description	anlee will be obing ad autoide
None retained	Local lab is being used, no san Lebanon, Lab samples are mai patient safety	nples will be shipped outside inly CBC, Chemistry to follow up c

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Target sample size 20	Actual enrollment target size 17
Date of first enrollment: Type Actual	Date of first enrollment: Date 10/08/2018
Date of study closure: Type Actual	Date of study closure: Date 31/12/2020
Recruitment status Complete	Recruitment status: Specify
Date of completion 12/01/2018	
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/NCT02941926? recrs=d&rslt=Without&type=Intr&cond=Advanced+Breast+Cancer&titles=co	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	
ΝΑ	



Contact for Public/Scientific Queries						
Contact type	Contact full name	Address	Country	Telephone	Email	Affiliation
Public	Fadi Farhat	Saida	Lebanon	+961 3 753 155	drfadi.trials@gm ail.com	Hammoud Hospital
Scientific	Hind Khairallah	Beirut	Lebanon	+961 1 512002 Ext. 271	Hind.Khairallah@ fattal.com.lb	Khalil Fattal et Fils s.a.l.
Public	Fadi El Karak	Mansourieh	Lebanon	+961 3 061 621	felkarak@yahoo. com	Bellevue Medical Center
Public	Georges Chahine	Beirut	Lebanon	+9613 647778	Chahine_george s@hotmail.com	Hotel Dieu 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	Georges Chahine	Chahine_georges@hotmail.com	009613 647778
Bellevue Medical Center	21/08/2017	Fadi El Karak	felkarak@yahoo.com	00961 3 061 621
Hammoud Hospital University Medical Center	02/05/2017	Fadi Farhat	drfadi.trials@gmail.com	00961 3 753 155



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, hemoglobin, platelets (as clinically indicated), Chemistry tests: Alkaline phosphatase, ALT (SGPT), AST (SGOT), calcium corrected for serum albumin, creatinine or creatinine clearance, potassium, sodium, magnesium,direct bilirubin, total bilirubin (as 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), , 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	