REPUBLIC OF LEBANON MINISTRY OF PUBLIC HEALTH Lebanon Clinical Trials Registry

Roll-over Study to Allow Continued Access to Ribociclib

11/09/2025 17:05:46

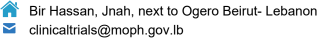
Main Information	
Primary registry identifying number	Protocol number
LBCTR2022095114	CLEE011A2412B
MOH registration number	
Study registered at the country of origin	Study registered at the country of origin: Specify
Yes	
Turne of unviolation	Tune of maniaturations, lundifie
Type of registration	Type of registration: Justify
Prospective	N/A
Date of registration in national regulatory	
agency	
Primary sponsor	Primary sponsor: Country of origin
Novartis Pharmaceuticals	Novartis Pharmaceuticals
Date of registration in primary registry	Date of registration in national regulatory agency
29/11/2022	
Dublic Alde	A
Public title	Acronym
Roll-over Study to Allow Continued Access to Ribociclib	
Scientific title	Acronym
A Post-trial Access Roll-over Study to Allow Access to Ribociclib	
(LEE011) for Patients Who Are on Ribociclib Treatment in Novartis- sponsored Study	
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	
ددة المراكز لتقتيم السلامة طويلة المدى لريبوسيكليب بالاشتراك مع أدوية أخرى في المشاركين الذين نوفارتيس، والتي أوفت بمتطلبات هدفها الأساسي (أهدافها)، ومن يعتقد طبيب الدراسة أنه سيستفيد من	هذه در اسه نمدید معوجه انتسمیه و متع یشار کون فی در اسهٔ عالمیهٔ تر عاها شرکهٔ ن
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 letro	ozole as was administered in their parent study)
Drug: Anastrozole (Participants continue ribociclib in combination with a Drug: Goserelin (Participants continue ribociclib in combination with gos	
Drug: Tamovifen (Participants continue ribociclib in combination with ta	

Drug: Fulvestrant (All participants continue ribociclib in combination with fulvestrant as was administered in their parent study)

Key inclusion and exclusion criteria: Inclusion criteria

REPUBLIC OF LEBANON MINISTRY OF PUBLIC HEALTH Lebanon Clinical Trials Registry

 Currently participating in a Novartis sponsored global study (parent study) and the parent study has fulfilled its primary objective(s) Must have been receiving treatment with ribociclib for at least 6 cycles in 13. Currently has evidence of clinical benefit as determined by the Investigated 	the parent study	o in combination with other drugs,
Key inclusion and exclusion criteria: Gender	Key inclusion and exclusion c	riteria: Specify gender
Both		
Key inclusion and exclusion criteria: Age minimum	Key inclusion and exclusion c	riteria: Age maximum
18	99	
Key inclusion and exclusion criteria: Exclusion criteria		
 Permanent discontinuation of ribociclib in the parent study Currently has unresolved toxicities for which ribociclib dosing has been in Local access to commercially available ribociclib and reimbursed 	terrupted in the parent study	
Type of study		
Interventional		
Type of intervention	Type of intervention: Specify t	ype
Pharmaceutical	N/A	
Trial scope	Trial scope: Specify scope	
Therapy	N/A	
	. .	
Study design: Allocation	Study design: Masking Open (masking not used)	
	Open (masking not used)	
Study design: Control	Study phase	
Uncontrolled	4	
Study design: Purpose	Study design: Specify purpose)
Treatment	N/A	
Study design: Assignment	Study design: Specify assignm	nent
Single	N/A	
IMP has market authorization	IMP has market authorization:	Specify
Yes, Lebanon and Worldwide	US, EU, and other countries	
Name of IMD	Veen of outbouingtion	Month of authorization
Name of IMP Ribociclib	Year of authorization 2017	1
	2017	
Type of IMP		
Others		
Pharmaceutical class		
highly selective small molecule inhibitor of cyclin-dependent kinases 4 and 6	6 (CDK4/6)	
Therapeutic indication		
Metastatic Breast Cancer		
Therapeutic benefit		
continued treatment to participants who are currently receiving ribociclib		
Study model	Study model: Explain model	
Study model N/A	Study model: Explain model	
1 1// 3		



REPUBLIC OF LEBANON Lebanon Clinical Trials Registry

Study model: Specify model N/A	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 None retained	Biospecimen description NA
Target sample size	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
Pending Date of completion	Recruitment status: Specify
-	Recruitment status: Specify 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.

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
Name

Secondary Sponsors Name NA

Contact for Public/Scientific Queries						
Contact type	Contact full name	Address	Country	Telephone	Email	Affiliation
Public	Jawad Makarem	Al Chouf	Lebanon	+961 3 484288	jawad.Makarem @awmedicalvilla ge.org	Ain Wazein Medical Village
Scientific	Hind Khairallah	Sin El Fil	Lebanon	+961 1 512002 Ext. 271	Hind.Khairallah@ fattal.com.lb	Khalil Fattal et Fils s.a.l
Public	Nagi El Saghir	Beirut	Lebanon	+961 3 827955	ns23@aub.edu.l b	American University of Beirut Medical Center

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





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 Kamaleddine	irb@awmedicalvillage.org	+961 (0) 5 509 001 ext 2014

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

REPUBLIC OF LEBANON Ministry of Public Health Lebanon Clinical Trials Registry

Health Conditions or Problems Studied		
Condition Code Keyword		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 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