

A Study of the Efficacy and Safety of Brolucizumab vs. Aflibercept in Patients With Visual Impairment Due to Diabetic Macular Edema

18/08/2025 13:09:20

Primary registry identifying number	Protocol number
LBCTR2019030200	RTH258B2302
MOH registration number	
31193/2018	
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 recently initiated, original file was previously submitted by Paper
Date of registration in national regulatory agency 23/07/2018	
Primary sponsor	Primary sponsor: Country of origin
Novartis Pharma Services Inc.	Novartis Pharmaceuticals
Date of registration in primary registry	Date of registration in national regulatory agency
15/12/2020	23/07/2018
Public title	Acronym
A Study of the Efficacy and Safety of Brolucizumab vs. Aflibercept in Patients With Visual Impairment Due to Diabetic Macular Edema	KITE
Scientific title	Acronym
A Two-Year, Two-Arm, Randomized, Double Masked, Multicenter, Phase III Study Assessing the Efficacy and Safety of Brolucizumab Versus Aflibercept in Adult Patients With Visual Impairment Due to Diabetic Macular Edema	
Brief summary of the study: English	
The purpose of this study is to evaluate the efficacy and safety of brolucizumab in treatment of patients with visual impairment due to diabetic macular edema (DME).	
Brief summary of the study: Arabic	
رجة التعمية، عشوائيّة التوزيع، من مجموعتين، مدّتها سنتان، لتقييم فعاليّة وسلامة دواء برولوسيزوماب مقابل دواء أفليبرسبت لدى المرضى البالغين المصابين بضعف بصريّ ناتج عن الوذمة البقعيّة السكريّ	در اسة مرحلة ثالثة، متعدّدة المراكز، مزدو
Health conditions/problem studied: Specify	
Patients With Visual Impairment Due to Diabetic Macular Edema	
Interventions: Specify	
•Drug: Brolucizumab Intravitreal injection	
Other Name: RTH258, ESBA1008	
•Drug: Aflibercept	

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Lebanon Clinical Trials Registry

Intravitreal injection		
Other Name: Eylea		
Key inclusion and exclusion criteria: Inclusion criteria		
 •Written informed consent before any assessment •Patients with type 1 or type 2 diabetes mellitus and HbA1c of ≤10% at scree •Medication for the management of diabetes stable within 3 months prior to of the study 		emain stable during the course
Key inclusion and exclusion criteria: Gender Both	Key inclusion and exclusion cr	riteria: Specify gender
Key inclusion and exclusion criteria: Age minimum	Key inclusion and exclusion cr	riteria: Age maximum
18	90	
Key inclusion and exclusion criteria: Exclusion criteria		
 Active proliferative diabetic retinopathy in the study eye Active intraocular or periocular infection or active intraocular inflammation in Uncontrolled glaucoma in the study eye defined as intraocular pressure (IO Previous treatment with anti-VEGF drugs or investigational drugs in the stu Stroke or myocardial infarction during the 6-month period prior to baseline Uncontrolled blood pressure defined as a systolic value ≥160 mmHg or diagonal 	P) > 25 millimeters mercury (mmH dy eye	g)
Other protocol-specified inclusion/exclusion criteria may apply		
Type of study		
Interventional		
Type of intervention	Type of intervention: Specify ty	уре
Pharmaceutical	N/A	
Trial scope Other	Trial scope: Specify scope	
Study design: Allocation	Study design: Masking	
Randomized controlled trial	Blinded (masking used)	
Study design: Control	Study phase	
Active	3	
Study design: Burnasa	Study design: Specify purpose	
Study design: Purpose Treatment	N/A	
Study design: Assignment Parallel	Study design: Specify assignm	nent
IMP has market authorization	IMP has market authorization:	Specify
NU		
Name of IMP	Year of authorization	Month of authorization
RTH258 (Brolucizumab)		
Type of IMP		
Immunological		
Pharmaceutical class		
Anti VEGF-A		
Therapeutic indication		



Diabetic Macular Edema Therapeutic benefit Change from baseline in best-corrected visual acuity (BCVA) at Week 52 Study model Study model: Explain model N/A Study model: Specify model **Time perspective** Time perspective: Explain time perspective N/A Time perspective: Specify perspective

Target follow-up duration

Number of groups/cohorts

Biospecimen retention Samples with DNA**

Target sample size

10

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N/A

N/A

N/A

N/A

Date of first enrollment: Type Actual

Date of study closure: Type Actual

Recruitment status Complete

Date of completion 19/06/2019

IPD sharing statement plan No

Biospecimen description

Target follow-up duration: Unit

Samples will be exported to : Q2 solutions The Alba campus Rosebank Livingston EH547EG United Kingdom Phone : 44 (0) 2033 184 884 x2401 Biosamples include Urine and Blood Urine for general analysis Blood : CBC, Chemistry, HbA1c, Lipids Panel, Anti Drug Ab, Pharmacogenomics

Actual enrollment target size

4

Date of first enrollment: Date 01/03/2019

Date of study closure: Date 22/12/2021

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

Additional data URL

https://clinicaltrials.gov/ct2/show/record/NCT03481660?term=CRTH258B2302&rank=1&view=record

Admin comments

Trial status

Approved

Secondary Identifying Numbers		
Full name of issuing authority	Secondary identifying number	
Clinical Trials. gov	NCT03481660	

Sources of Monetary or Material Support	
Name	
Novartis Pharma Services Inc.	

Secondary Sponsors

Name

NA

Contac	Contact for Public/Scientific Queries					
Contact type	Contact full name	Address	Country	Telephone	Email	Affiliation
Public	Naji Waked	Beirut	Lebanon	009613252 552	wakednaji@yaho o.com	Hotel Dieu De France
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	Joseph Nehme	Dbayeh	Lebanon	009610313 6120	drjosephnehme @gmail.com	Eye and Ear Hospital Internation al
Public	Hala El Rami	Beirut	Lebanon	76367510	ramielhala@hot mail.com	Beirut Eye and ENT specialist Hospital

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Centers/Hospitals Involved in the Study			
Conter/Hospital name I Name of principles investidator		Principles investigator speciality	Ethical approval
Hotel Dieu De France	Naji Waked	Ophthalmology	Approved
Eye and Ear Hospital International	Joseph Nehme	Ophthalmology	Approved
Beirut Eye and ENT specialist Hospital	Hala El Rami	Ophthalmology	Approved

Ethics Review	Ethics Review			
Ethics approval obtained	Approval date	Contact name	Contact email	Contact phone
Hotel Dieu de France	30/04/2018	Sami Richa	cue@usj.edu.lb	961421229
Other Hotel Dieu De France (Eye and Ear Hospital International)	02/10/2018	Sami Richa	cue@usj.edu.lb	961421229
Beirut Eye and ENT Specialist Hospital	21/12/2018	Sami Richa	cue@usj.edu.lb	961421229



Countries of Recruitment
Name
Lebanon
Belgium
Bulgaria
Czech Republic
Denmark
Estonia
France
Germany
Hungary
India
Republic of Korea
Latvia
Lithuania
Malaysia
Norway
Singapore
Slovakia
Sweden
Switzerland
Turkey

Health Conditions or Problems Studied		
Condition Code Keyword		Keyword
Diabetic macular edema	Oedema, unspecified (R60.9)	Macular Edema





Interventions		
Intervention	Description	Keyword
Physical Exam, Vital signs, ophtalmic Exam, IOP, Optical Coherence Tomography, Fluorescein Angiography, Color Fundus photography, Urinalysis, Serum/ urine pregnancy test, lab test, completion of QoL questionnaires	ICF, Lab, questionnaires, Medication administration, physical examination	ICF, Lab tests, Questionnaires, Medication administration

Primary Outcomes		
Name	Time Points	Measure
Change from baseline in best-corrected visual acuity (BCVA)	Baseline, week 52	baseline, week 52

Key Secondary Outcomes		
Name	Time Points	Measure
Average change from baseline in BCVA	wk 40 till wk 52	wk 40 till wk 52
Proportion of patients with injections per planned dosing regimen	wk8,12,16	wk8,12,16
Change from baseline in central subfield thickness	baseline up to wk 100	baseline up to wk 100



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