

Open-Label Extension of Voxelotor

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Main Information

Primary registry identifying number

LBCTR2020063513

MOH registration number

Study registered at the country of origin

Yes

Type of registration

Prospective

Date of registration in national regulatory agency

25/06/2020

Primary sponsor

Global Blood Therapeutics Inc.

Date of registration in primary registry

26/10/2022

Public title

Open-Label Extension of Voxelotor

Scientific title

An Open-Label Extension Study of Voxelotor Administered Orally to Participants with Sickle Cell Disease Who Have Participated in

Voxelotor Clinical Trials

Brief summary of the study: English

Open-label extension (OLE) study of Voxelotor for participants with Sickle Cell Disease who have participated in Voxelotor clinical trials. Approximately 600 participants with sickle cell disease (SCD), aged ≥ 4 to >18year years will be enrolled at approximately 70 global clinical sites. Participants aged ≥ 12 years will receive a voxelotor dose of 1500 mg QD, regardless of their body weight. Participants aged < 12 years will receive a voxelotor dose based on

their body weight, to provide exposure corresponding to the adult dose of 1500 mg QD. The participant's weight at study entry will be used to determine the starting voxelotor dose in this study. The dose should be adjusted if the participant's weight increases or decreases at a scheduled clinic visit. The objective of this OLE is to assess the safety of, and SCD-related complications of, long-term treatment with Voxelotor, in participants who have completed treatment in a Global Blood Therapeutics (GBT)-sponsored Voxelotor clinical study.

Brief summary of the study: Arabic

أعوام و المصابين بمرض الخلايا المنجلية و النين شاركوا في التجارب السريرية للالمشاركين ابتداءاً من عمر voxelotor دراسة تكميلية على voxelotor السابقة على مستحضر

Health conditions/problem studied: Specify

Sickle Cell Disease

Protocol number

GBT440-038

Study registered at the country of origin: Specify

Type of registration: Justify

N/A

Primary sponsor: Country of origin

United States of America

Date of registration in national regulatory agency

25/06/2020

Acronym

Acronym



Interventions: Specify

Drug: Voxelotor (GBT440)

All participants will receive voxelotor once daily (QD), administered orally as tablets, dispersible tablets, or powder for oral suspension formulation

Key inclusion and exclusion criteria: Inclusion criteria

1. Male or female participant with SCD, aged ≥ 4 years, who participated and received study drug in a GBT-sponsored voxelotor pediatric clinical study

Note: Participants who discontinued study drug due to an AE, but who remained on study, may be eligible for treatment in this study provided the AE does not pose a risk for treatment with voxelotor.

2. Female participants of childbearing potential are required to have a negative urine pregnancy test before dosing on Day 1.

Note: Female participants who become childbearing during the study must be willing to have a negative urine pregnancy test to remain in the study

- 3. If sexually active, female participants of childbearing potential must use highly effective methods of contraception until 30 days after the last dose of study drug. If sexually active, male participants must use barrier methods of contraception until 30 days after the last dose of study drug.
- 4. Participant has provided written assent (both the consent of the participant's legal representative or legal guardian and the participant's assent

[where applicable] must be obtained)

5. Subjects with abnormal TCD who have not completed Study GBT440-032 can participate in OLE study

Key inclusion and exclusion criteria: Gender Key inclusion and exclusion criteria: Specify gender

Both

Key inclusion and exclusion criteria: Age minimum Key inclusion and exclusion criteria: Age maximum

Key inclusion and exclusion criteria: Exclusion criteria

- 1. Female participant who is breastfeeding or pregnant
- 2. Participant withdrew consent from a GBT-sponsored voxelotor pediatric clinical study
- 3. Participant was lost to follow-up from a GBT-sponsored voxelotor pediatric clinical study
- 4. Participant has any medical, psychological, safety, or behavioral conditions that, in the opinion of the investigator, may confound safety interpretation.

interfere with compliance, or preclude informed consent

5. Based on the most recent Oxbryta® US label (December 2021), co-administration with both moderate and strong CYP3A4 inducers should be avoided

Type of study

Interventional

Type of intervention Type of intervention: Specify type

Pharmaceutical

Trial scope Trial scope: Specify scope

Therapy

Study design: Allocation Study design: Masking

N/A Open (masking not used)

Study design: Control Study phase

Active 3

Study design: Purpose Study design: Specify purpose

Treatment

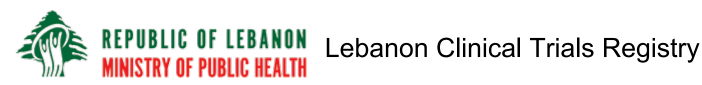
Study design: Assignment Study design: Specify assignment

Single

IMP has market authorization IMP has market authorization: Specify

Yes, Worldwide United States of America

Name of IMP Year of authorization Month of authorization



Voxelotor (Oxbryta) 2019 11

Type of IMP

Others

Pharmaceutical class

Allosteric modulator of hemoglobin oxygen affinity

Therapeutic indication

Sickle Cell Disease

Therapeutic benefit

Voxelotor is an orally bioavailable HbS polymerization inhibitor that binds specifically to HbS with a 1:1 stoichiometry, and exhibits preferential partitioning to RBCs. By increasing Hb's affinity for oxygen, voxelotor inhibits HbS polymerization in a dose dependent manner that may improve deformability, decrease the viscosity of SCD blood, and ultimately increase blood flow in the microcirculation, thus improving net O2 delivery. Therefore, chronically modifying 20% to 30% of HbS with voxelotor in subjects with SCD is expected to deliver the clinical benefits of reducing HbS polymerization while improving O2 delivery to peripheral tissues.

Study model: Explain model Study model

N/A N/A

Study model: Specify model

N/A

Time perspective Time perspective: Explain time perspective

Time perspective: Specify perspective

Target follow-up duration Target follow-up duration: Unit

Number of groups/cohorts

Biospecimen retention Biospecimen description

None retained N/A

Target sample size Actual enrollment target size

21

Date of first enrollment: Type Date of first enrollment: Date

11/08/2020 Anticipated

Date of study closure: Type Date of study closure: Date

30/06/2026 Anticipated



24



Recruitment status Pending	Recruitment status: Specify
Date of completion	
IPD sharing statement plan No	IPD sharing statement description N/A
Additional data URL	
Admin comments	
Trial status Approved	

Secondary Identifying Numbers		
Full name of issuing authority	Secondary identifying number	
Clinicaltrials.gov	NCT04188509	
WHO International Clinical Trials Registry Platform	EUCTR2019-003144-76-GB	

Sources of Monetary or Material Support	
Name	
Global Blood Therapeutics Inc. USA	

Secondary Sponsors		
Name		
N/A		



Contac	Contact for Public/Scientific Queries					
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Public	Dr. Adlette Inati	Tripoli	Lebanon	961322803 3	adlette.inati@lau. edu.lb	Nini Hospital

Centers/Hospitals Involved in the Study			
Center/Hospital name Name of principles investigator		Principles investigator speciality	Ethical approval
American University of Beirut Medical Center	Dr. Miguel Abboud	Hematology	Not approved
Nini Hospital	Dr. Adlette Inati	Hematology	Approved

Ethics Review	Ethics Review				
Ethics approval obtained	Approval date	Contact name	Contact email	Contact phone	
Nini Hospital	15/06/2020	Dr. Nabil Kabbara	N/A	9616431400	



Countries of Recruitment
Name
United States of America
United Kingdom
Lebanon
Brazil
Egypt
Oman
Kenya
Nigeria
Ghana

Health Conditions or Problems Studied		
Condition	Code	Keyword
Sickle-Cell Disorder	Sickle-cell disorders (D57)	Hematology, Sickle Cell, Disorder

Interventions			
Intervention	Description	Keyword	
Drug	Voxelotor	GBT440	

Primary Outcomes			
Name	Time Points	Measure	
Treatment Emergent Adverse Events and Serious Adverse Events	Throughout entire study	N/A	
Sickle Cell Disease-Related Complications	Throughout entire study	Frequency of SCD-related complications	

Key Secondary Outcomes		
Name	Measure	
N/A	N/A	N/A



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	