**REPUBLIC OF LEBANON** MINISTRY OF PUBLIC HEALTH Lebanon Clinical Trials Registry

# Study to Evaluate the Effect of GBT440 in Pediatrics With Sickle Cell Disease

18/07/2025 02:12:30

Primary registry identifying number	Protocol number
BCTR2019090195	GBT440-007
1011 as a interaction an unable of	
AOH registration number 288/م	
مر/2001	
Study registered at the country of origin	Study registered at the country of origin: Specify
/es	
ype of registration	Type of registration: Justify
Retrospective	Requested by Sponsor- Registry not in place upon study initiation
Date of registration in national regulatory	
agency 1/06/2014	
Primary sponsor	Primary sponsor: Country of origin
Global Blood Therapeutics Inc.	United States of America
Date of registration in primary registry 30/09/2019	Date of registration in national regulatory agency 11/06/2014
0/03/2013	11/00/2014
Public title	Acronym
Study to Evaluate the Effect of GBT440 in Pediatrics With Sickle Cell Disease	
Scientific title	Acronym
A Phase 2a, Open-label, Single and Multiple Dose Study to Evaluate the Pharmacokinetics, Safety, Tolerability, and Exploratory Freatment Effect of GBT440 in Pediatric Participants With Sickle Cell Disease	
Brief summary of the study: English	
This study consists of three parts, Parts A, B, and C. Part A is a single dose PK study in pediatric participants with Sickle Cell Disease. (Closed on 07 Aug 2017 (LPLV)) Part B is a multiple dose, safety, exploratory, efficacy, and PK study in adolescent Sickle Cell Disease participants who were 12- 17 years of age (Closed on 04 Jan 2019 (LPLV)) Part C is a multiple dose, safety, tolerability, and PK study, which ncludes the assessment of hematological effects and the effect on ICD flow velocity of GBT440 is pediatric participants with Sickle Cell Disease who are 4 to 17 years of age.	
Brief summary of the study: Arabic	
در اسة لتقييم تأثير GBT440 عند الأطغال المصابين بمرض الخلايا المنجلي	
lealth conditions/problem studied: Specify	
Sickle Cell Disease	

Drug: GBT440 administered as oral capsules, tablets or dispersible tablets



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Key inclusion and exclusion criteria: Inclusion criteria		
<ol> <li>Male or female participants with homozygous hemoglobin SS (HbSS)</li> <li>Age:         <ul> <li>Part A – 6 to 17 years of age. (Cohort 1 [12 to 17] and Cohort 2 [6 to 11] as defined in the Study Design)</li> <li>Part B – 12 to 17 years of age</li> <li>Part C – 4 to 17 years of age</li> <li>A participant taking hydroxyurea (HU) may be enrolled if the dose has</li> </ul> </li> </ol>	-	
<ul> <li>adjustments during the study and no sign of hematological toxicity.</li> <li>4. Hemoglobin (Hb):</li> <li>Part A – No restriction</li> <li>Part B – Hb ≤10.5 g/dL</li> <li>Part C – Hb ≤10.5 g/dL</li> <li>5. Written informed parental/guardian consent and participant assent ha (EC) policy and requirements, consistent with ICH guidelines.</li> </ul>		
Key inclusion and exclusion criteria: Gender	Key inclusion and exclusion	criteria: Specify gender
Both	,	
Key inclusion and evolution evidence. And minimum		
Key inclusion and exclusion criteria: Age minimum 4	Key inclusion and exclusion	i chiena: Age maximum
4	17	
Key inclusion and exclusion criteria: Exclusion criteria		
<ol> <li>Any one of the following requiring medical attention within 14 days pri</li> <li>Vaso-occlusive crisis (VOC)</li> <li>Acute chest syndrome (ACS)</li> <li>Splenic sequestration crisis</li> <li>Dactylitis</li> <li>Requires chronic transfusion therapy</li> </ol>		
<ol> <li>History of stroke or meeting criteria for primary stroke prophylaxis (his</li> <li>Transfusion within 30 days prior to signing the ICF</li> </ol>	tory of two TCD measurements ≥20	00 cm/sec)
Type of study		
Interventional		
Type of intervention	Type of intervention: Specify	y 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	
Dose comparison	2	
Study design: Purpose	Study design: Specify purpo	222
Treatment	N/A	56
Study design: Assignment	Study design: Specify assig	nmont
Single	N/A	linent
Single	14/74	
IMP has market authorization No	IMP has market authorization	n: Specify
Name of IMP	Year of authorization	Month of authorization
GBT440 (Voxelotor)		
Type of IMP Others		



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Ρ	ha	rma	ceu	tical	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	Study model: Explain model
N/A	N/A
Study model: Specify model	
N/A	
Time perspective	Time perspective: Explain time perspective
N/A	N/A
Time perspective: Specify perspective	
N/A	
Target follow-up duration	Target follow-up duration: Unit

Number of groups/cohorts

Biospecimen retention
None retained

Biospecimen description N/A

**Target sample size** 24

Date of first enrollment: Type
Actual

Date of study closure: Type Actual

Recruitment status Recruiting

Date of completion

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Actual enrollment target size 32

Date of first enrollment: Date 21/07/2016

Date of study closure: Date 31/01/2021

**Recruitment status: Specify** 



# IPD sharing statement plan IPD sharing statement description No N/A Additional data URL Yata and the second se

Secondary Identifying Numbers		
Full name of issuing authority	Secondary identifying number	
Clinicaltrials.gov	NCT02850406	
EU Clinical Trials Registry	EudraCT: 2016-004209-15	

### Sources of Monetary or Material Support

### Name

Global Blood Therapeutics, Inc. USA

### **Secondary Sponsors**

No Sponsors



Contac	Contact for Public/Scientific Queries					
Contact type	Contact full name	Address	Country	Telephone	Email	Affiliation
Public	Dr. Adlette Inati	Tripoli	Lebanon	961322803 3	adlette.inati@lau. edu.lb	Nini Hospital
Scientific	Dan Rudin	171 Oyster Point Blvd., Suite 300 South San Francisco, CA 94080	United States of America	(650) 534- 2574	drudin@gbt.com	Global Blood Therapeuti cs
Public	Dr. Miguel Abboud	Beirut	Lebanon	961135000 0	ma56@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	Dr. Miguel Abboud	Pediatric Hematology- Oncology	Approved
Rafik Hariri University Hospital	Dr. Adlette Inati	Pediatric Hematology- Oncology	Approved
Nini Hospital	Dr. Adlette Inati	Pediatric Hematology- Oncology	Approved

Ethics Review				
Ethics approval obtained	Approval date	Contact name	Contact email	Contact phone
American University of Beirut Medical Center	09/07/2018	Dr. Fuad Ziyadeh	irb@aub.edu.lb	9611738025
Rafic Hariri University Hospital	31/08/2018	Dr. Iyad Issa	NA	9611830000
Nini Hospital	31/08/2018	Dr. Nabil Kabbara	NA	9616431400

### **Countries of Recruitment**

Name
Lebanon
United States of America
United Kingdom



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Health Conditions or Problems Studied		
Condition Code Keyword		Keyword
Sickle Cell Disease	Sickle-cell disorders (D57)	Sickle Cell, Anemia, Hemolytic, Congenital, Hematologic Diseases

Interventions		
Intervention	Description	Keyword
Drug	GBT440	Oral Capsule, Tablet or Dispersible Tablet

Primary Outcomes				
Name	Time Points	Measure		
Part A: Pharmacokinetic profile of GBT440 including maximum concentration	Pre-dose to Day 15	Pharmacokinetic profile		
Part A: Pharmacokinetic profile of GBT440 including the time taken to reach the maximum concentration	Pre-dose to Day 15	Pharmacokinetic profile		
Part A: Pharmacokinetic profile of GBT440 including the total drug concentration over time	Pre-dose to Day 15	Pharmacokinetic profile		
Part B: Change in hemoglobin	Baseline to Week 24	Hemoglobin in Blood		
Part C: Change in cerebral blood flow	Baseline to Week 48	TAMM TCD velocity		

### **Key Secondary Outcomes**

Name	Time Points	Measure	
Part A: Number of participants with treatment-related adverse events	Days 1 - 15	Assessed by CTCAE v4.03	
Part B: Multiple Dose effect on Clinical Measures of Hemolysis	Day 1 - Week 24	Clinical Measures of Hemolysis	
Part B: Pharmacokinetic profile of GBT440 including maximum concentration	Pre-dose to Week 24	Pharmacokinetic profile	
Part B: Pharmacokinetic profile of GBT440 including the time taken to reach the maximum concentration	Pre-dose to Week 24	Pharmacokinetic profile	
Part B: Pharmacokinetic profile of GBT440 including the total drug concentration over time	Pre-dose to Week 24	Pharmacokinetic profile	
Part C: Multiple dose effect on clinical measures of hemolysis	Baseline to Week 24 and Week 48	Clinical Measures of Hemolysis	
Part C: Change in cerebral blood flow	Baseline to Week 24	Measured by the TAMM TCD velocity	
Part C: Pharmacokinetic profile of GBT440 including maximum concentration	Pre-Dose to Week 48	Pharmacokinetic profile	
Part C: Pharmacokinetic profile of GBT440 including the time taken to reach the maximum concentration	Pre-Dose to Week 48	Pharmacokinetic profile	
Part C: Pharmacokinetic profile of GBT440 including the total drug concentration over time	Pre-Dose to Week 48	Pharmacokinetic profile	



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	

