

## An Open Label Extension Study of PTG-300 in Non-Transfusion Dependent (NTD) and Transfusion-Dependent (TD) β-Thalassemia Subjects

12/08/2025 18:20:06

_				4 8		
IN.	n	•	~~	41	$\sim$	-
ı.						

Primary registry identifying number

LBCTR2019070220

MOH registration number

Study registered at the country of origin

Type of registration

Prospective

Date of registration in national regulatory

Date of registration in primary registry

An Open Label Extension Study of PTG-300 in Non-Transfusion Dependent (NTD) and Transfusion-Dependent (TD) β-Thalassemia

Dependent (NTD) and Transfusion-Dependent (TD) β-Thalassemia

This is an open-label, long term extension study for subjects completing study PTG-300-02. After completing the previous study, eligible subjects who choose to continue treatment may enroll in the PTG-300-03 study. The safety evaluation done at the end of study PTG-300-02 will be used to confirm subject eligibility for this study (see Screening in Study Procedure section). No interruption of PTG

19/06/2019

**Primary sponsor** 

Protagonist Therapeutics Inc

08/07/2019

**Public title** 

Subjects

Scientific title An Open Label Extension Study of PTG-300 in Non-Transfusion

Subjects Brief summary of the study: English

-300 treatment is expected with the transition between studies. Brief summary of the study: Arabic

سيتم استخدام تقييم السلامة الذي تم في نهاية الدراسة PTG-300-02 Health conditions/problem studied: Specify

دراسة تمديدية مفتوحة التسمية للأشخاص الذين اكملوا دراسة 02

Chronic anemia due to ineffective erythropoiesis (IE) in subjects with  $\beta$  thalassemia

-PTG-300. بعد الانتهاء من الدراسة السابقة, قد يتم تسجيل الاشخاص المؤهلين الذين يختارون مواصلة العلاج في دراسة PTG-300-03

لتاكيد اهلية الموضوع لهذه الدراسة لا يتوقع اي انقطاع للعلاج مع الانتقال بين الدراسات.

Interventions: Specify

Protocol number

PTG300-03

Study registered at the country of origin: Specify

Type of registration: Justify

N/A

Primary sponsor: Country of origin

Date of registration in national regulatory agency

19/06/2019

Acronym

Acronym

Bir Hassan, Jnah, next to Ogero Beirut- Lebanon clinicaltrials@moph.gov.lb



Subjects rolling over from the PTG-300-02 study, who meet the response criteria defined for this study at the last dose received in study PTG-300-02, will continue to receive the same dose in PTG-300-03.

Subjects who did not meet response criteria defined for this study at the last dose received in study PTG-300-02, will start PTG-300-03 at the next higher dose level.

#### Key inclusion and exclusion criteria: Inclusion criteria

- 1. NTD and TD β-thalassemia subjects who completed Week 12 and Week 16 respectively in Study PTG-300-02.
- 2. Women of childbearing potential (WOCBP) and men agree to use a highly effective contraceptive measure (base on the Clinical Trial Facilitation

Group [CTFG]) during the duration of the study and for 4 weeks after the last dose of study drug in the case of women and 90 days after the

dose of study drug in the case of men.

- 3. For WOCBP, a negative urine pregnancy test within 24 hours prior to the first dose of study medication in this study.
- 4. Subjects or legal guardians (in the case of minors) understand the study procedures and agree to participate in the study by giving written informed consent.
- 5. Subjects, or legal representative (in the case of minors) are willing and able to adhere to the study visit schedule and other protocol requirements.
- 6. Subjects between 12-<18 years of age understand and provide the assent to participate in the study, according to local guidelines.

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. Subjects who discontinued prematurely from study 300-02 (before Week 12 in NTD and Week 16 in TD)
- 2. Clinically meaningful laboratory abnormalities at Screening.
- 3. Pregnant or lactating females.
- 4. Current history of alcohol dependence or illicit drug use.
- 5. Subject has a concurrent clinically significant, unstable or uncontrolled cardiovascular, pulmonary, hepatic, renal, gastrointestinal, genitourinary.

hematological, coagulation, immunological, endocrine/metabolic or other medical disorder that, in the opinion of the Investigator, might confound the results of the study or pose additional risk to the subject by their participation in the study.

- 6. Subject is mentally or legally incapacitated at the time of Screening visit or has a history of clinically significant psychiatric disorders that would impact the subject's ability to participate in the trial according to the Investigator.
- 7. Concurrent participation in any other interventional study.

### Type of study

Interventional

N/A

Type of intervention Type of intervention: Specify type

Pharmaceutical N/A

Trial scope Trial scope: Specify scope

Therapy

Study design: Allocation Study design: Masking N/A: Single arm study Open (masking not used)

Study phase Study design: Control

Study design: Purpose Study design: Specify purpose

Treatment

Study design: Assignment Study design: Specify assignment

N/A Single

IMP has market authorization IMP has market authorization: Specify

Year of authorization Name of IMP Month of authorization



PTG-300

### Type of IMP

Cell therapy

#### Pharmaceutical class

PTG-300 is a peptidic agent structurally related to natural hepcidin that mimics its inhibitory activity on

### Therapeutic indication

β thalassemia

### Therapeutic benefit

Administration of PTG-300 may result in iron redistribution in  $\beta$ -thalassemia subjects with potentially beneficial effects on erythropoiesis and consequently improvements in chronic anemia. This improvement in ineffective erythropoiesis may result in a clinical benefit both in NTD and in TD  $\beta$ thalassemia subjects, by improving the symptomatology of the chronic anemia and the complications of the extramedullary hematopoiesis in the first group and by decreasing the need for transfusions in the latter.

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

Time perspective: Specify perspective

N/A

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

Number of groups/cohorts

Biospecimen retention

Samples without DNA

Biospecimen description

Blood samples taken throughout the study will be shipped to ICON lab in Ireland for analysis. These samples will be then stored at ICON Lab by Protagonist Therapeutics for up to 10 vears.

Target sample size

84

Date of first enrollment: Type

Anticipated

Date of study closure: Type

Anticipated

Actual enrollment target size

Date of first enrollment: Date

05/08/2019

Date of study closure: Date

27/06/2022



Recruitment status

Pending

Date of completion

25/06/2020

IPD sharing statement plan

No

**Recruitment status: Specify** 

IPD sharing statement description

Not applicable

Additional data URL

none

Admin comments

**Trial status** 

Approved

Secondary Identifying Numbers	
Full name of issuing authority	Secondary identifying number
Food and Drug Administration	IND

## **Sources of Monetary or Material Support**

Name

Protagonist Therapeutics.inc

### **Secondary Sponsors**

Name

Not Applicable

Contac	Contact for Public/Scientific Queries						
Contact type	Contact full name	Address	Country	Telephone	Email	Affiliation	
Public	Aziz Zoghbi	MCT-CRO, Berytech Technology and Health, 5th Floor Damascus Road, Beirut, Lebanon	Lebanon	009611612 500	zog_az@mct- cro.com	Regional Manager	
Scientific	Ali Taher	Chronic Care Center, Hazmieh, Lebanon	Lebanon	009613755 669	ataher@aub.edu. lb	PI	



Centers/Hospitals Involved in the Study				
Center/Hospital name	Name of principles investigator	Principles investigator speciality	Ethical approval	
Chronic Care Center	Dr. Ali Taher	Hematology/Oncology	Approved	

Ethics Review					
Ethics approval obtained	Approval date	Contact name	Contact email	Contact phone	
Chronic Care Center	01/06/2019	Michele Abi Saad	cccmas@chroniccare.org.lb	05-455101	

Countries of Recruitment
Name
Lebanon
Thailand
United Kingdom
United States of America
Turkey
Tunisia
Malaysia
Greece
Italy

Health Conditions or Problems Studied		
Condition	Code	Keyword
Thalassemia	Thalassaemia (D56)	Thalassemia



Interventions				
Intervention	Description	Keyword		
PTG-300	3mg/week	NA		
PTG-300	10mg/week	NA		
PTG-300	20mg/week	NA		
PTG-300	40mg/week	NA		
PTG-300	40 mg every 2 weeks	NA		
PTG-300	80 mg/week	NA		
PTG-300	80 mg every 2 weeks	NA		

Primary Outcomes				
Name	Time Points	Measure		
NTD Subjects who achieve an increase in Hgb ≥1.0 g/dL	from pre-treatment baseline without transfusion	Hemoglobin Test		
NTD patients: Hgb change	from pre-treatment baseline	Hemoglobin Test		
TD Subjects who achieve a ≥20% reduction in the red blood cell (RBC) units transfused over 8-week period compared to pre-treatment baseline	over an 8-week period	RBC units transfused		
TD patients: Change in the number of units of RBC required	from pre-treatment baseline	RBC units transfused		



Key Secondary Outcomes				
Name	Time Points	Measure		
NTD patients: Proportion of subjects who achieve an increase in Hgb ≥1.5 g/dL	from pre-treatment baseline without transfusion	Hgb test		
Proportion of subjects who achieve a maintenance dose	from pre-treatment baseline	Dose		
Hgb level	from pre-treatment baseline	Hgb test		
NTD Patients: Duration of Hgb change of ≥1.0 g/dL	from pre-treatment baseline without transfusion	Hgb test		
NTD Patients: Duration of Hgb change of ≥1.5 g/dL from	from pre-treatment baseline without transfusion	Hgb test		
Change in the following PD parameters	from pre-treatment baseline	serum iron, ferritin, transferrin saturation (TSAT)		
TD Patients: Proportion of subjects who achieve ≥ 33% reduction in the RBC units required over an 8-week period	over an 8-week period	RBC units transfused		
TD Patients: Duration of response defined as ≥ 20% reduction in the RBC units	over an 8-week period	RBC units transfused		
TD Patients: Number of RBC units required	from pre-treatment baseline	RBC units transfused		
TD Patients: Percent change in the RBC units required	from pre-treatment baseline	RBC units transfused		
TD: Hgb change	from pre-treatment baseline	Hgb test		

7



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	