

Study registered at the country of origin: Specify

Date of registration in national regulatory agency

LCTR was recently initiated, original file was previously submitted

Safety Study of Crushed Deferasirox Film Coated Tablets in Pediatric Patients With Transfusional Hemosiderosis (MIMAS)

Protocol number

Type of registration: Justify

Primary sponsor: Country of origin

Novartis Pharmaceuticals

CICL670F2429

by Paper

01/08/2018

Acronym

Acronym

06/07/2025 17:47:04

Main Information

Primary registry identifying number

LBCTR2019030206

MOH registration number

32772/2018

Study registered at the country of origin

Type of registration

Retrospective

Date of registration in national regulatory

01/08/2018

Primary sponsor

Novartis Pharma Services Inc.

Date of registration in primary registry

08/07/2019

Public title

Safety Study of Crushed Deferasirox Film Coated Tablets in Pediatric Patients With Transfusional Hemosiderosis (MIMAS)

Scientific title

A Single-arm Interventional Phase IV, Post-authorisation Study Evaluating the Safety of Pediatric Patients With Transfusional Hemosiderosis Treated With Deferasirox Crushed Film Coated

Brief summary of the study: English

The study employs an interventional, prospective, single arm, open label, global, multicenter, non-randomized trial design to monitor and assess the safety profile of the crushed deferasirox FCT in pediatric patients between age ≥2 to <6 with transfusional hemosiderosis over 24 weeks. This study will aim to enroll at least 40 patients.

Brief summary of the study: Arabic

دراسة تدخلية وحيدة المجموعة في المرحلة الرابعة بعد الترخيص لتقييم سلامة المرضى الاطفال المصابين بالحدد الدموي (الهيموسيدروسز) ذي الصلة بنقل الدم والمعالجين بأقراص ديفيرازيروكس المسحوقة المغلفة بطبقة رقيقة

Health conditions/problem studied: Specify

Transfusional Hemosiderosis

Interventions: Specify

Drug: Deferasirox

Deferosirox is provided in tablet forms of 90, 180 and 360mg. Tablets must be crushed.

Other Name: ICL670

Key inclusion and exclusion criteria: Inclusion criteria

1.Patients ≥2 to <6 years old diagnosed with transfusional hemosiderosis



- 2.Documented history of red blood cell transfusions
- 3. Written informed consent/assent before any study-specific procedures. The consent will be obtained from caregiver(s) or patient's legal representative. Investigators will also obtain assent of patients according to local, regional, or national regulations.
- 4.For patients on prior DFX: Serum ferritin (SF) >500 ng/mL, measured at screening visit 1 and requiring a DFX daily dose equivalent to FCT ≥
- 5.For patients on a prior chelator other than DFX (e.g. deferiprone or deferoxamine) or chelation naive: Serum ferritin (SF) >1000 ng/mL measured at screening visits 1 and 2.

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

Roth

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

Key inclusion and exclusion criteria: Exclusion criteria

- 1.Patients that receive more than one iron chelator at the same time as current iron chelation treatment. (Patients who have received combination therapy in their medical history but are currently being treated with a single ICT agent are eligible.)
- 2.Patients continuing on deferoxamine or deferiprone in addition to study treatment.

(Patients switching to or continuing on deferasirox are eligible).

- 3.Unresolved adverse events if the patient was previously treated with deferiprone or deferoxamine or deferasirox.
- 4. Significant proteinuria as indicated by a urinary protein/creatinine ratio > 0.5 mg/mg in a non-first void sample urine measured at screening
- 5. Serum creatinine > age adjusted ULN measured at any screening visit
- 6.Creatinine clearance below 90 mL/minute measured at any screening visit. Creatinine clearance using the Schwartz formula will be estimated from serum creatinine measured at each respective visit.
- 7.ALT and/or AST > 2.5 x ULN measured at screening visit 1.
- 8.Total bilirubin (TBIL) >1.5 x ULN measured at screening visit 1.
- 9.Patients with significant impaired GI function or GI disease that may significantly alter the absorption of oral deferasirox FCT (e.g. ulcerative diseases, uncontrolled nausea, vomiting, diarrhea, malabsorption syndrome, or small bowel resection).
- 10. History of and/or laboratory evidence of active Hepatitis B or Hepatitis C (HBsAg in the absence of HBsAb OR HCV Ab positive with HCV RNA positive.
- 11.Liver disease with severity of Child-Pugh Class B or C.
- 12. History of hypersensitivity to any of the study drug or excipients.
- 13. Patients participating in another clinical trial or receiving an investigational drug.
- 14. Patients with a known history of HIV seropositivity.
- 15. Patients unwilling or unable to comply with the protocol.
- 16. History of malignancy of any organ system, treated or untreated, within the past 5 years whether or not there is evidence of local recurrence or metastases, with the exception of localized basal cell carcinoma of the skin.
- 17. Significant medical condition interfering with the ability to partake in this study (e.g.

uncontrolled hypertension, unstable cardiac disease not controlled by standard medical therapy, systemic disease: cardiovascular, renal, hepatic, etc.).

18. Female patients who reach menarche and they or their caregivers refuse pregnancy testing and/or if there is a positive pregnancy test

Type of study

Interventional

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

Single

IMP has market authorization

Yes, Lebanon and Worldwide

Name of IMP

Deferasirox Film Coated Tablets "Jadenu"

Type of IMP

Others

Pharmaceutical class

Non-chiral, Tridentate ligand iron chelator

Therapeutic indication

Iron Over Load

Therapeutic benefit

Treatment of Iron Overload symptoms

Study model

N/A

Study model: Specify model

N/A

Time perspective

N/A

Time perspective: Specify perspective

N/A

Target follow-up duration

Number of groups/cohorts

Biospecimen retention

None retained

Target sample size

3

Date of first enrollment: Type

N/A

Study design: Specify assignment

IMP has market authorization: Specify

USA, UK, France, Germany, Netherlands, Switzerland, Sweden,

Year of authorization

Month of authorization

2017 10

Study model: Explain model

N/A

Time perspective: Explain time perspective

Target follow-up duration: Unit

Biospecimen description

Local Lab tests at site

Actual enrollment target size

3

Date of first enrollment: Date





Actual	10/10/2018

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

Actual 11/03/2020

Recruitment status Recruitment status: Specify

Recruiting

Date of completion

22/04/2019

IPD sharing statement plan IPD sharing statement description

No undecided

Additional data URL

https://clinicaltrials.gov/ct2/show/record/NCT03372083?id=CICL670F2429&rank=1&view=record

Admin comments

Trial status

Approved

Secondary Identifying Numbers	
Full name of issuing authority	Secondary identifying number
Clinical Trials. gov	NCT03372083

Sources of Monetary or Material Support

Name

Novartis Pharma Services Inc.

Secondar	v Sponsors

Name

NA



Contac	Contact for Public/Scientific Queries					
Contact type	Contact full name	Address	Country	Telephone	Email	Affiliation
Public	Ali Taher	Beirut	Lebanon	009613755 669	ataher@aub.edu. lb	Chronic Care Center
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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	Approved

Ethics Review				
Ethics approval obtained	Approval date	Contact name	Contact email	Contact phone
Chronic Care Center	26/07/2018	Michele Abi saad	cccmas@chroniccare.org.lb	+961 3 664 310
American University of Beirut Medical Center	05/07/2018	Fuad Ziyadeh	fz05@aub.edu.lb	+961 (0) 1 350 000 ext:5445

Countries of Recruitment
Name
Egypt
Lebanon
Italy
Oman
Russian Federation
Saudi Arabia
Thailand
United Arab Emirates



Health Conditions or Problems Studied		
Condition Code Keyword		
Transfusional Hemosiderosis	Thalassaemia, unspecified (D56.9)	Transfusional Hemosiderosis

Interventions			
Intervention	Description	Keyword	
Physical examination, Vitals, Height, Weight, Hematology, Chemistry, urinalisis, ECG, Ocular assessment, Auditory assessment	Physical examination, Vitals, Height, Weight, Hematology, Chemistry, urinalisis, ECG, Ocular assessment, Auditory assessment	ICF, IMP, Lab tests , diary completion	

Primary Outcomes			
Name	Time Points	Measure	
Percentage of patients with selected gastrointestinal disorders	24 weeks	24 wks	
To assess the safety of crushed deferasirox FCT with respect to selected gastrointestinal (GI) disorders	through out the study	through out the study	

Key Secondary Outcomes			
Name	Time Points	Measure	
Percentage of patients who experienced AEs suspected to be related to study drug	24 weeks	24 wks	
•Change from baseline ECGs up	24 weeks	24 weeks	
•Change from baseline serum ferritin (SF)	24 weeks	24 weeks	
Absolute change for serum creatinine	24 weeks	24 weeks	
•Absolute change for creatinine clearance UPCR	24 weeks	24 weeks	
Palatability Questionnaire Score	24 weeks	24 weeks	



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	