REPUBLIC OF LEBANON MINISTRY OF PUBLIC HEALTH Lebanon Clinical Trials Registry

Long Term Extension Study in Patients With Primary Hyperoxaluria

20/08/2025 05:53:45

Main Information	
Primary registry identifying number LBCTR2020124677	Protocol number DCR-PHXC-301
MOH registration number NCT04042402	
Study registered at the country of origin No	Study registered at the country of origin: Specify Study registered in clinicaltrials.gov
Type of registration Prospective	Type of registration: Justify N/A
Date of registration in national regulatory agency 02/08/2019	
Primary sponsor Dicerna Pharmaceuticals, Inc.	Primary sponsor: Country of origin USA
Date of registration in primary registry	Date of registration in national regulatory agency
20/02/2021	02/08/2019
Public title Long Term Extension Study in Patients With Primary Hyperoxaluria	Acronym
Scientific title An Open-Label Roll-Over Study to Evaluate the Long-Term Safety and Efficacy of DCR-PHXC Solution for Injection (Subcutaneous	Acronym "PHYOX3"
Use) in Patients With Primary Hyperoxaluria	
The proposed study is designed to provide patients previously enrolled in Phase 1 and 2 studies of DCR-PHXC long-term access to DCR-PHXC, and to evaluate the long-term safety and efficacy of DCR-PHXC in patients with PH.	
Brief summary of the study: Arabic	
ا تم تصميم الدراسة المقترحة لتزويد المرضى المسجلين سابقًا في دراسات المرحلتين الأولى والثانية من PH. على المدى الطويل في مرضى DCR-PHXC ولتقييم سلامة وفعالية ، DCR-PHXC إلى	بالوصول طويل الأمد DCR-PHXC
Health conditions/problem studied: Specify	
Primary Hyperoxaluria Type 1 (PH1) Primary Hyperoxaluria Type 2 (PH2) Kidney Diseases Urologic Diseases Genetic Disease	
Interventions: Specify	
Drug: DCR-PHXC Multiple fixed doses of DCR-PHXC by subcutaneous (SC) injection. Other Name: Nedosiran	
Key inclusion and exclusion criteria: Inclusion criteria	
Key Inclusion Criteria:	

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•Participant successfully completed a Dicerna Pharmaceuticals, Inc. study of DCR PHXC.

younger than 18 years of age and must have genetically confirmed PH.

•For participants rolling over from a multidose study of DCR-PHXC, enrollment should occur within a window of 25 to 60 days from the last dose of study intervention. Estimated GFR at screening ≥ 30 mL/min normalized to 1.73 m2 body surface area (BSA), calculated using Chronic Kidney Disease Epidemiology Collaboration (CKD EPI) formula in participants aged ≥ 18 years (Levey & Stevens, 2010), or the formula by Schwartz in participants aged 6 to 16 years (Schwartz et al., 2009; National Kidney Foundation, 2002). In Japan, the formula by Uemura et al. will be used for participants aged 6 to 17 years (Uemura et al., 2014). 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 6 99 Key inclusion and exclusion criteria: Exclusion criteria Kev exclusion criteria include: · Renal or hepatic transplantation; prior or planned within the study period · Current dialysis · Documented evidence of clinical manifestations of systemic oxalosis (including pre-existing retinal, heart, or skin calcifications, or history of severe bone pain, pathological fractures, or bone deformations). Type of study Interventional Type of intervention Type of intervention: Specify type Pharmaceutical N/A **Trial scope** Trial scope: Specify scope Other Study design: Allocation Study design: Masking Single Arm Study Open (masking not used) Study design: Control Study phase N/A 3 Study design: Purpose Study design: Specify purpose Treatment N/A Study design: Assignment Study design: Specify assignment Single N/A IMP has market authorization IMP has market authorization: Specify No Name of IMP Year of authorization Month of authorization DCR PHXC Type of IMP Others

OR Participant is the sibling of a participant who successfully completed a Dicerna Pharmaceuticals, Inc. study of DCR PHXC. Siblings must be

Pharmaceutical class

A synthetic double-stranded (hybridized duplex) ribonucleic acid (RNA) oligonucleotide conjugated to N-acetyl-D-galactosamine (GalNAc) amino-sugar residues.

Therapeutic indication

Primary Hyperoxaluria.



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Therapeutic benefit At present, no therapies are approved by regulatory authorities for the treatm DCR-PHXC treatment has the potential benefit to reduce or eliminate the ex- the liver and thus avoid the need for a combined liver and kidney transplanta already on renal replacement therapy.	nent of patients with PH. cess oxalate production in tion in patients not
Study model	Study model: Explain model
N/A	N/A
Study model: Specify model N/A	
Time perspective N/A	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	Biospecimen description
None retained	Blood and Urine Samples
Target sample size 50	Actual enrollment target size
Date of first enrollment: Type	Date of first enrollment: Date
Anticipated	01/02/2021
Date of study closure: Type	Date of study closure: Date
Anticipated	30/12/2023
Recruitment status	Recruitment status: Specify
Other	Enrolling by Invitation
Date of completion 30/12/2023	
IPD sharing statement plan	IPD sharing statement description
No	Participants will be assigned a unique identifier by the Sponsor. Any participant records or datasets that are transferred to the Sponsor will contain the identifier only; participant names or any information which would make the participant identifiable will not

be transferred.

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Additional	data	URL
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Admin comments

Trial status

Approved

Secondary Identifying Numbers		
Full name of issuing authority	Secondary identifying number	
US NCT Number	NCT04042402	

Sources of Monetary or Material Support

Name

Dicerna pharmaceuticals inc. 75 Hayden Avenue Suite 400 Lexington, MA 02421, USA

Secondary Sponsors

Name

NA

Contact for Public/Scientific Queries						
Contact type	Contact full name	Address	Country	Telephone	Email	Affiliation
Public	Chadi Safa	lebanon. Baabda. Chiah. Ain el remeneh	Lebanon	009617125 1819	Chadi.safa@clin art.net	Clinart
Scientific	Chebl Mourani	Alfred Naccache Blvd, External Viewing Tower, Floor 4, Room 9403	Lebanon	03 290090	cheblmourani@g mail.com	HDF

Centers/Hospitals Involved in the Study			
Center/Hospital name	Name of principles investigator	Principles investigator speciality	Ethical approval
1.Hotel Dieu de France	Chebl Mourani	Pediatric Nephrologist	Approved
2.Saint George University Hospital	Pauline Abu jaoude	Nephrologist	Approved





Ethics Review				
Ethics approval obtained	Approval date	Contact name	Contact email	Contact phone
Hotel Dieu de France	30/03/2020	Nancy Choukair Alam	nancy.alam@usj.edu.lb	: +961 1 421 000
Saint George Hospital University Medical Center	09/07/2020	Sandra Berberi	smberbari@stgeorgehospital.org	+961 1 44 16 30

Countries of Recruitment

Name
France
Netherlands
Germany
United Kingdom
United States of America
Lebanon
Spain
Italy
Australia
Canada
Japan

Health Conditions or Problems Studied			
Condition	Code	Keyword	
Primary Hyperoxaluria	2-Propanol (T51.2)	Kidney Diseases	

Interventions		
Intervention	Description	Keyword
DCR-PHXC	Multiple fixed doses of DCR-PHXC by subcutaneous (SC) injection	Nedosiran





Primary Outcomes			
Name	Time Points	Measure	
To evaluate the effect of DCR PHXC on estimated glomerular filtration rate	Annual change from baseline	estimated glomerular filtration rate	

Key Secondary Outcomes		
Name	Time Points	Measure
The incidence and severity of treatment-emergent adverse events (TEAE) and SAEs associated with abnormal 12 lead electrocardiogram (ECG) readings	TEAEs and SAEs are evaluated monthly for 3 years	Electrocardiogram (ECG)

Date of first journal publication of results

Trial Results

Summary results

Study results globally

Date of posting of results summaries

Results URL link

Baseline characteristics

Participant flow

Adverse events

Outcome measures

URL to protocol files

