REPUBLIC OF LEBANON Lebanon Clinical Trials Registry MINISTRY OF PUBLIC HEALTH

Phase 2 Placebo-Controlled, Double-Blind, Multicenter Study to Evaluate the Efficacy, Safety, and Tolerability of DCR-PHXC Solution for Injection (subcutaneous use) in Patients with Primary Hyperoxaluria

20/08/2025 13:21:05 Main Information Primary registry identifying number Protocol number LBCTR2020043435 DCR-PHXC-201 MOH registration number Study registered at the country of origin Study registered at the country of origin: Specify Study registered in clinicaltrials.gov Type of registration Type of registration: Justify Prospective N/A Date of registration in national regulatory agency 15/02/2019 **Primary sponsor** Primary sponsor: Country of origin Dicerna Pharmaceuticals, Inc US Date of registration in primary registry Date of registration in national regulatory agency 27/04/2020 15/02/2019 Public title Acronym Phase 2 Placebo-Controlled, Double-Blind, Multicenter Study to Evaluate the Efficacy, Safety, and Tolerability of DCR-PHXC Solution for Injection (subcutaneous use) in Patients with Primary Hyperoxaluria Scientific title Acronym PHYOX2 Phase 2 Placebo-Controlled, Double-Blind, Multicenter Study to Evaluate the Efficacy, Safety, and Tolerability of DCR-PHXC Solution for Injection (subcutaneous use) in Patients with Primary Hyperoxaluria Brief summary of the study: English This is a 6-month randomized, placebo-controlled, double-blind study of DCR-PHXC in patients with primary hyperoxaluria (PH1 and PH2). Potential participants are screened over an up-to- 6week period prior to randomization to DCR-PHXC or placebo. The proposed study is designed to evaluate the efficacy, safety, tolerability, and PK of DCR-PHXC versus placebo in patients with PH1 and PH2.

Brief summary of the study: Arabic

-DCR أشهر ، في المرحلة الثانية، متعددة المراكز ومزدوجة التعمية، محكومة بدواء وهمي لتقييم فعالية و سلامة محلول الحقن6دراسة مدتها (تحت الجلد) ودرجة تحمله لدى المرضى الذين يعانون من فرط أوكسالات البول الأولي PHXC

Health conditions/problem studied: Specify

DCR-PHXC is designed to selectively reduce LDHA messenger ribonucleic acid (mRNA) and lactate dehydrogenase (LDH) activity in the liver, and subsequently decrease liver oxalate production. DCR-PHXC is being developed as a treatment for PH, an ultra-rare autosomal recessive



No

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disease characterized by excessive production of oxalate in the liver.

Interventions: Specify

DCR-PHXC is a synthetic RNAi drug that consists of a double-stranded oligonucleotide conjugated to GalNAc ligands. DCR-PHXC is a sterile formulation of drug substance (DCR-L1360) in WFI, intended for SC administration. DCR-PHXC is not commercially available in any country. The placebo comparator is 0.9% normal saline for injection.

Key inclusion and exclusion criteria: Inclusion criteria

• 24-hour Uox excretion ≥ 0.7 mmol (adjusted per 1.73 m2 body surface area [BSA] in participants < 18 years of age) in both collections performed in the screening period. Of the first 24 participants enrolled, at least 12 (50%) must have at least one 24-hour Uox excretion ≥ 1.6 mmol (adjusted per 1.73 m2 BSA in participants aged < 18 years).

• Less than 20% variation between the two 24-hour urinary creatinine excretion values [mmol/24 hr/kg] derived from the two 24-hour urine collections in the screening period.

• Estimated glomerular filtration rate (eGFR) at screening ≥ 30 mL/min normalized to 1.73 m2 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 17 years, (Schwartz et al., 2009; National Kidney Foundation, 2002). In Japan, the formula by Uemura et al. will be used for

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participants aged 6 to 17 years.

| 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

• Renal or hepatic transplantation (prior or planned within the study period)

· Current dialysis or anticipated requirement for dialysis during the study period

Plasma oxalate > 30 µmol/L

• Documented evidence of clinical manifestations of systemic oxalosis (including preexisting retinal, heart, or skin calcifications, or history of severe bone pain, pathological fractures, or bone deformations)

• Liver function test (LFT) abnormalities: Alanine aminotransferase (ALT) and/or aspartate aminotransferase (AST) >1.5 times upper limit of normal (ULN) for age and gender.

Type of study

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Interventional

| Type of intervention Pharmaceutical | Type of intervention: Specify type N/A | |
|---|--|------|
| Trial scope Other | Trial scope: Specify scope | |
| Study design: Allocation Randomized controlled trial | Study design: Masking Blinded (masking used) | |
| Study design: Control Placebo | Study phase 2 | |
| Study design: Purpose Treatment | Study design: Specify purpose N/A | |
| Study design: Assignment Parallel | Study design: Specify assignment N/A | |
| IMP has market authorization No | IMP has market authorization: Specify | |
| Name of IMP DCR-PHXC | Year of authorizationMonth of authoriza20201 | tion |
| Type of IMP | | |



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Study model: Explain model

Others

Pharmaceutical class

Synthetic double-stranded (hybridized duplex) RNA oligonucleotide conjugated to GalNAc .

Therapeutic indication

DCR-PHXC is designed to selectively reduce LDHA messenger ribonucleic acid (mRNA) and lactate dehydrogenase (LDH) activity in the liver, and subsequently decrease liver oxalate production. DCR-PHXC is being developed as a treatment for PH, an ultra-rare autosomal recessive disease characterized by excessive production of oxalate in the liver.

Therapeutic benefit

Study model

Patients with PH are predisposed to the development of multiple and recurrent urinary tract (urolithiasis) and kidney (nephrolithiasis) stones. This deposition of calcium oxalate in the renal parenchyma produces tubular toxicity and renal damage. At present, no therapies are approved by regulatory authorities for the treatment of patients with PH. A number of supportive therapies are used in an attempt to mitigate some of the effects of the disease. DCR-PHXC treatment has the potential benefit to reduce or eliminate the excess oxalate production in

the liver and thus avoid the need for a combined liver and kidney transplantation in patients not already on renal replacement therapy.

| N/A | N/A |
|---|---|
| Study model: Specify model N/A | |
| Time perspective N/A Time perspective: Specify perspective N/A | Time perspective: Explain time perspective N/A |
| Target follow-up duration | Target follow-up duration: Unit |
| Number of groups/cohorts | |
| Biospecimen retention | Biospecimen description |
| Samples with DNA** | Samples may be stored for a maximum of 5years (or according to local regulations) following the last participant's last visit for the study at a facility selected by the sponsor to enable further analysis of immune responses to DCR-PHXC. |
| Target sample size 36 | Actual enrollment target size |
| Date of first enrollment: Type Actual | Date of first enrollment: Date 28/10/2019 |
| Date of study closure: Type Actual | Date of study closure: Date 30/01/2021 |

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| Recruitment status | Recruitment status: Specify |
|----------------------------|---|
| Recruiting | |
| Date of completion | |
| IPD sharing statement plan | IPD sharing statement description |
| No | Participants will be assigned a unique identifier by the Sponsor. Any participant records or data sets 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 |
| Additional data URL | |

Admin comments

Trial status

Approved

| Secondary Identifying Numbers | | | |
|--------------------------------|------------------------------|--|--|
| Full name of issuing authority | Secondary identifying number | | |
| 1.US NCT | NCT03847909 | | |
| 2.WHO Universal | U1111-1224-6881 | | |

Sources of Monetary or Material Support

Name

Dicerna pharmaceuticals inc. 87 Cambridgepark Drive Cambridge, MA 02140 US

| Secondary Sponsors | |
|--------------------|--|
| Name | |
| N/A | |

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| Contact for Public/Scientific Queries | | | | | | |
|---------------------------------------|-------------------|---------|---------|-------------------|----------------------------|-------------------------|
| Contact type | Contact full name | Address | Country | Telephone | Email | Affiliation |
| Public | Chadi Safa | Beirut | Lebanon | 009617125 1819 | chadi.safa@clina rt.net | Clinart Mena |
| Scientific | Chebl Mourani | Beirut | Lebanon | 009611290 090 | cheblmourani@g mail.com | Hotel Dieu de France |

| Centers/Hospitals Involved in the Study | | | | |
|---|---------------------------------|------------------------------------|------------------|--|
| Center/Hospital name | Name of principles investigator | Principles investigator speciality | Ethical approval | |
| Hotel Dieu De France | Chebl Mourani | Pediatric Nephrology | Approved | |
| Saint George University Hospital | Pauline Abou Jaoude | Pediatric Nephrology | Approved | |

| Ethics Review | | | | |
|---|---------------|----------------|--------------------------------|---------------|
| Ethics approval obtained | Approval date | Contact name | Contact email | Contact phone |
| Hotel Dieu de France | 01/10/2019 | Nancy Alam | nancy.alam@usj.edu.lb | 01421000 |
| Saint George Hospital University Medical Center | 29/11/2019 | Sandra Berberi | smberbari@stgeorgehospital.org | 01 1 44 16 30 |





| Countries of Recruitment |
|--------------------------|
| Name |
| Australia |
| Canada |
| France |
| Germany |
| Italy |
| Japan |
| Netherlands |
| New Zealand |
| Poland |
| Romania |
| Spain |
| United Kingdom |
| United States of America |
| Lebanon |

| Health Conditions or Problems Studied | | | |
|---------------------------------------|-----------------------------------|-------|--|
| Condition Code Keyword | | | |
| Primary Hyperoxaluria | Nephrotic syndrome, other (N04.8) | РНҮОХ | |

| Interventions | | | |
|---------------|--|----------|--|
| Intervention | Description | Keyword | |
| DCR-PHXC | DCR-PHXC is a synthetic RNAi drug that consists of a double-stranded oligonucleotide conjugated to GalNAc ligands. DCR-PHXC is a sterile formulation of drug substance (DCR- L1360) in WFI, intended for SC administration. DCR-PHXC is not commercially available in any country. | DCR-PHXC | |
| Placebo | The placebo comparator is 0.9% normal saline for injection. | Placebo | |





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| Primary Outcomes | | | |
|---|--|-------------|--|
| Name | Time Points | Measure | |
| To assess the efficacy of DCR-PHXC in reducing urinary oxalate burden in patients with PH (types 1 and 2) | The proportion of participants with a reduction from baseline in 24-hour Uox of at least 70%, based on a TWS AUC and/or reaching normalization or near- normalization of 24-hour Uox on at least 2 consecutive visits, starting from Day 90. Normalization of Uox is defined as < 0.46 mmol/24 hours; near- normalization is defined as ≥ 0.46 to < 0.60 mmol/24 hours (values adjusted per 1.73 m2 BSA in participants aged < 18 years). | 24-hour Uox | |

| Key Secondary Outcomes | | | |
|--|--|---|--|
| Name | Time Points | Measure | |
| To evaluate the effect of DCR-PHXC on stone burden in patients with PH | Percent change in the summed surface area and number of kidney stones identified via kidney ultrasound from Baseline to Day 180 | Number of Kidney stone | |
| To evaluate the effect of DCR-PHXC on plasma oxalate in patients with PH | Percent change in plasma oxalate from Baseline to Day 180 (for adults only) | Plasma Oxalate | |
| To evaluate the effect of DCR-PHXC on eGFR | Rate of change in eGFR from Baseline to Day 180 | eGFR | |
| To assess the safety of DCR-PHXC in patients with PH | AE and SAE; change from Baseline in 12-lead ECG, physical examination findings, vital signs, and clinical laboratory tests | 12 Lead ECG, Physical Examination test, Vital signs, Clinical Laboratory test. | |
| To characterize the PK of DCR-PHXC in patients with PH | Population and individual PK parameters for DCR- PHXC | РК | |





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