

## CLCZ696B2319E1 Open Label Extension Study to Evaluate Long-term Safety of Sacubitril/Valsartan in Pediatric Patients With Heart Failure (HF).

12/09/2025 08:31:14

| Main Information   |  |
|--|--|
| Primary registry identifying number  | Protocol number                                    |
| LBCTR2019070266  | CLCZ696B2319E1                                     |
| MOH registration number  |  |
| Study registered at the country of origin  | Study registered at the country of origin: Specify |
| Yes  |  |
| Type of registration   | Type of registration: Justify                      |
| Prospective  | N/A  |
| Date of registration in national regulatory agency   |  |
| Primary sponsor  | Primary sponsor: Country of origin                 |
| Novartis Pharma Services   | Novartis Pharmaceuticals                           |
| Date of registration in primary registry   | Date of registration in national regulatory agency |
| 11/04/2022   |  |
| Public title   | Acronym  |
| CLCZ696B2319E1 Open Label Extension Study to Evaluate Long-<br>term Safety of Sacubitril/Valsartan in Pediatric Patients With Heart<br>Failure (HF).   |  |
| Scientific title   | Acronym  |
| A multicenter study to evaluate long-term safety and tolerability of<br>open label sacubitril/valsartan in pediatric patients with heart failure<br>due to systemic left ventricle systolic dysfunction who have<br>completed study CLCZ696B2319 |  |
| Brief summary of the study: English  |  |
| The purpose of this study is to evaluate long-term safety and<br>tolerability data in eligible CLCZ696B2319 (PANORAMA-HF)<br>patients receiving open-label<br>sacubitril/valsartan   |  |
| Brief summary of the study: Arabic   |  |
| ة الأمد لدواء ساكيوبيتريل / فالسارتان المفتوح اللصاقة وقدرة تحمّله لدى أطفال مرضى مصابين بغشل<br>القلب بسبب الخلل الوظيفي الانقباضي الجهازي للبُطيُّن الأيسر وقد<br>الجزوا دراسة CLCZ696B2319  | در اسة متعددة المراكز لتقييم السلامة الطويل        |
| Health conditions/problem studied: Specify   |  |
| Heart failure patients   |  |
| Interventions: Specify   |  |
| Drug: sacubitril/valsartan   |  |
| Target dose 3.1 mg/kg bid  |  |

 $\sim$ 



#### Formulations:

1

Tablets (50, 100, 200 mg) Granules [12.5 mg (4 granules), 31.25 mg (10 granules), in capsules] Liquid (1 mg/ml, 4 mg/ml, prepared from tablets)

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

Signed informed consent

Other Name: LCZ696

On study drug at PANORAMA-HF Part 2 End of Study visit. Does not have any significant safety issue

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

 Both
 Revenue of the second second

Key inclusion and exclusion criteria: Age minimum

Key inclusion and exclusion criteria: Age maximum 18

#### Key inclusion and exclusion criteria: Exclusion criteria

Subject only participated in PANORAMA-HF Part 1 or was a Screen Failure in PANORAMA-HF or permanently discontinued study drug in PANORMA-HF Part 2

Use of investigational drugs within 5 half-lives of enrollment or within 30 days (longer duration); with the exception of PANORAMA-HF study drug (requires >/=36-hour washout before baseline visit)

History of hypersensitivity or allergy to study treatment, its excipients or drugs of similar chemical class, ACEIs, ARBs, or NEP inhibitor and known/suspected contraindications to sacubitril/valsartan

Renal vascular hypertension (including renal artery stenosis)

Significant renal estimated glomerular filtration rate disorder (eGFR calculated using modified Schwartz formula <30% mean GFR for age); hepatic disorder (serum aspartate aminotransferase or alanine aminotransferase > 3 times upper limit of normal); gastrointestinal disorder or biliary disorder

History of angioedema

Parents or legal guardians of subject who do not give consent or allow the child to give assent, or inability of patient or parents/legal guardians to follow instructions or comply with follow-up procedures

Any medical condition(s) that may put the patient at risk in the investigator's opinion or that the investigator deems unsuitable for the study Other protocol defined inclusion/exclusion criteria may apply

#### Type of study

Interventional

| Type of intervention   | Type of intervention: Specify ty  | vpe   |
|--|---|---|
| Pharmaceutical   | N/A   |   |
| Trial scope  | Trial scope: Specify scope  |   |
| Safety   | N/A   |   |
| Study design: Allocation   | Study design: Masking   |   |
| N/A: Single arm study  | Open (masking not used)   |   |
| Study design: Control  | Study phase   |   |
| N/A  | 2   |   |
|  |   |   |
| Study design: Purpose  | Study design: Specify purpose   |   |
| Study design: Purpose<br>Treatment                                     | Study design: Specify purpose N/A   |   |
|  |   |   |
| Treatment  | N/A   |   |
| Treatment Study design: Assignment                                     | N/A<br>Study design: Specify assignm  | ent   |
| Treatment Single   | N/A<br>Study design: Specify assignm<br>N/A   | ent<br>Specify<br>the dosage forms 50,100 and |
| Treatment Study design: Assignment Single IMP has market authorization | N/A<br>Study design: Specify assignm<br>N/A<br>IMP has market authorization: S<br>Lebanon and worldwide : yes for t | ent<br>Specify<br>the dosage forms 50,100 and |



# Lebanon Clinical Trials Registry

| Type of IMP   |  |
|---|--|
| Others  |  |
| Pharmaceutical class  |  |
| LCZ696, also known as Entresto® (sacubitril/valsartan) is an angiotensin rec<br>(ARNI), providing concomitant neprilysin (neutral endopeptidase 24.11, NEP<br>angiotensin II type 1 (AT1) receptor blockade |  |
| Therapeutic indication  |  |
| Pediatric patients with heart failure   |  |
| Therapeutic benefit   |  |
| long-term safety and tolerability data in eligible CLCZ696B2319 (PANORAM open-label sacubitril/valsartan.   | A-HF) patients receiving                   |
| 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<br>N/A  |  |
|   |  |
| Target follow-up duration   | Target follow-up duration: Unit            |
| Number of groups/cohorts  |  |
|   |  |
| Biospecimen retention   | Biospecimen description                    |
| Samples without DNA   | blood and urine samples                    |
|   |  |
|   |  |
|   |  |
|   |  |
| Target sample size  | Actual enrollment target size              |
| 10  | 9  |
| Date of first enrollment: Type  | Date of first enrollment: Date             |
| Actual  | 03/01/2020                                 |
| Date of study closure: Type   | Date of study closure: Date                |
| Actual  | 30/12/2022                                 |
| Recruitment status  | Pocruitment status: Specify                |
| Complete  | Recruitment status: Specify                |
|   |  |
| Date of completion 13/12/2021   |  |





| IPD sharing statement plan<br>No   | IPD sharing statement description Undecided |
|--|---|
|  |   |
| Additional data URL<br>https://clinicaltrials.gov/ct2/show/record/NCT03785405?cond=pediatric+hea | rt+failure&rank=8&view=record               |
| Admin comments   |   |
| <b>Trial status</b><br>Approved  |   |

| Secondary Identifying Numbers  |                              |
|--------------------------------|------------------------------|
| Full name of issuing authority | Secondary identifying number |
| Clinicaltrials.gov             | NCT03785405                  |

| Sources of Monetary or Material Support |
|---|
| Name                                    |
| Novartis Pharma Services                |

| Secondary Sponsors |  |
|--------------------|--|
| Name               |  |
| NA                 |  |

| Contac          | t for Public/Scientific Queries | 5          |         |                              |                                   |  |
|-----------------|---------------------------------|------------|---------|------------------------------|-----------------------------------|--|
| Contact<br>type | Contact full name               | Address    | Country | Telephone                    | Email                             | Affiliation                            |
| Public          | Linda Daou                      | Beirut     | Lebanon | 961604976                    | drlindadaou@gm<br>ail.com         | Hotel Dieu                             |
| Scientific      | Hind Khairallah                 | Sin El Fil | Lebanon | +961 1<br>512002<br>Ext. 271 | Hind.Khairallah@<br>fattal.com.lb | Khalil<br>Fattal et<br>Fils s.a.l.     |
| Public          | Ghassan Chehab                  | Beirut     | Lebanon | 961338858<br>1               | ghassanchehab<br>@yahoo.com       | Rafik Hariri<br>UNiversity<br>Hospital |





| Centers/Hospitals Involved in the Study |                                 |                                    |                  |
|---|---------------------------------|------------------------------------|------------------|
| Center/Hospital name                    | Name of principles investigator | Principles investigator speciality | Ethical approval |
| Hotel Dieu de France                    | Linda Daou                      | Pediatric Cardiology               | Approved         |
| Rafik Hariri University Hospital        | Ghassan Chehab                  | Cardiology                         | Approved         |

| Ethics Review                       |               |              |                          |                    |
|-------------------------------------|---------------|--------------|--------------------------|--------------------|
| Ethics approval obtained            | Approval date | Contact name | Contact email            | Contact phone      |
| Hotel Dieu de France                | 06/06/2019    | Sami Richa   | cue@usj.edu.lb           | 961421229          |
| Rafic Hariri University<br>Hospital | 28/01/2019    | Rawan Yamout | rawan.yamout@crurhuh.com | 018300000 ext 2037 |

### **Countries of Recruitment**

| Name           |
|----------------|
| Austria        |
| Argentina      |
| Canada         |
| Croatia        |
| Czech Republic |
| Egypt          |
| Finland        |
| France         |
| Germany        |
| Hungary        |
| India          |
| Japan          |
| Jordan         |
| Poland         |
|                |



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

| Portugal                            |
|-------------------------------------|
| Romania                             |
| Russian Federation                  |
| Saudi Arabia                        |
| Singapore                           |
| South Africa                        |
| Democratic People Republic of Korea |
| Spain                               |
| Sweden                              |
| Switzerland                         |
| Thailand                            |
| Turkey                              |
| United Kingdom                      |
| United States of America            |
| Lebanon                             |

| Health Conditions or Problems Studied |                     |         |  |
|---------------------------------------|---------------------|---------|--|
| Condition                             | Code                | Keyword |  |
| Heart Failure                         | Heart failure (I50) | HF      |  |

| Interventions                      |                                    |                                    |
|------------------------------------|------------------------------------|------------------------------------|
| Intervention                       | Description                        | Keyword                            |
| ICF, Physical Exam, ECG, Lab tests | ICF, Physical Exam, ECG, Lab tests | ICF, Physical Exam, ECG, Lab tests |





| Primary Outcomes   |                                   |                         |  |  |
|--|-----------------------------------|-------------------------|--|--|
| Name   | Time Points                       | Measure                 |  |  |
| Number of participants with Adverse Events (AEs) as a measure of safety and tolerability           | to end of study, up to 3<br>years | safety and tolerability |  |  |
| .Number of participants with Serious Adverse Events (SAEs) as a measure of safety and tolerability | to end of study, up to 3<br>years | safety and tolerability |  |  |

| Name                    | Time Points | Measure |
|-------------------------|-------------|---------|
| No secondary objectives | NA          | NA      |

#### **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                |  |
|                                      |  |