## **ALP**ORT SYNDROME

EFFICACY & SAFETY TRIAL-1

VONAFEXOR is an investigational drug

already tested in **nine studies** in patients with **other diseases** 

Orphan Drug

Designation for

Alport syndrome



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preserve kidney function and

reverse kidney damage in patients with Alport syndrome

## Participant's eligibility



The inclusion criteria for the study are:

- 16\*−55 years of age, woman or man
- Alport diagnosis history & genetic test results positive for Alport syndrome
- Mild to moderate loss of kidney function
- Increased albuminuria
- ACE, ARB or SGLT2i treatment allowed\*\*

## **About ALPESTRIA-1**

- This Phase 2 study will evaluate:
  - 1. the **safety** of Vonafexor in Alport syndrome
  - 2. the benefit of three dose levels of Vonafexor on renal function and biomarkers.
- The study aims at enrolling 20 patients in US and EU (France, Germany and Spain)
- All participants will receive Vonafexor as oral tablets once daily for 24 weeks then stop the treatment for 12 weeks in the follow-up period.
  No placebo all subjects will receive study drug.
- Assessments are a mix of 5 site visits, 4 home visits and 4 phone calls, with urine and/or blood sampling





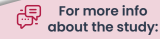


Vonafexor treatment

12 weeks
No Vonafexor treatment
(follow-up)

\* 16 years of age for the US ; 18 years of age for the EU countries

\*\* ACEi: angiotensin converting enzyme Inhibitor ARB: angiotensin receptor blocker SGLT2i: sodium-glucose cotransporter-2 inhibitor



Please visit: www.alportsyndrome.org

Once the study is registered: www.clinicaltrials.gov and www.clinicaltrialsregister.eu



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