



A Phase 2, Open-Label, Basket Study of Atrasentan in Patients with Proteinuric Glomerular Diseases

Status: Recruiting

Eligibility Criteria

Sex: Male or Female
Age Group: Not specified

This study is NOT accepting healthy

volunteers

Inclusion Criteria:

- Age 18 years and older for patients in the IgAN, FSGS, and Alport Syndrome cohorts - age 18-70 years for patients in the DKD cohort - receiving a maximally tolerated dose of RAS inhibitor therapy (ACEi or ARB) that has been stable for at least 12 weeks - there are different requirements for each diagnosis category & study staff will review these

Exclusion Criteria:

- current diagnosis of another cause of chronic kidney disease or another primary glomerulopathy - history of kidney transplantation or other organ transplantation - except for FSGS patients, use of systemic immunosuppressant medications, such as steroids, for more than 2 weeks in the past 3 months - blood pressure above 150 mmHg systolic or 95 mmHg diastolic - history of heart failure or a previous hospital admission for fluid overload. - history of liver disease - hemoglobin below 9 g/dL or blood transfusion for anemia within the past 3 months. - cancer in the past 5 years (except nonmelanoma skin cancer and curatively treated cervical carcinoma in situ) - women who are pregnant, breastfeeding, or intend become pregnant during the study - recently received an investigational agent -clinically significant unstable or uncontrolled medical condition (study staff will review)

Conditions & Interventions

Conditions:

Kidney, Prostate & Urinary

Keywords:

Glomerular Disease, Alport Syndrome, IgAN, FSGS, Proteinuric Glomerular Diseases

More Information

Description: The purpose of the research is to find out if atrasentan delays worsening of kidney function in IgAN, FSGS, and Alport Syndrome.

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Phase: Phase 2

IRR

Number: STUDY00012146

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