AFFINITY: A Phase 2 Basket Trial to Study the Safety and Efficacy of Atrasentan in Multiple Proteinuric Diseases

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Chinook is a clinical-stage biopharmaceutical company discovering, developing and commercializing precision medicines for rare, severe chronic kidney diseases.
Chinook’s Commitment to Kidney Disease Drug Development

Advancing pipeline of precision medicines for kidney diseases

<table>
<thead>
<tr>
<th>Program</th>
<th>Indication</th>
<th>Target Validation</th>
<th>Lead Optimization</th>
<th>IND-Enabling</th>
<th>Phase 1</th>
<th>Phase 2</th>
<th>Phase 3</th>
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<tbody>
<tr>
<td>Atrasentan</td>
<td>IgA Nephropathy</td>
<td>Phase 3 enrollment commenced in early 2021</td>
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<td>Basket of glomerular diseases</td>
<td>Phase 2 enrollment commenced in early 2021</td>
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<td>BION-1301</td>
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<td>Research Programs</td>
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<td>Discovery Programs</td>
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We continue to evaluate opportunities to add additional kidney disease programs to pipeline
Modernization of Drug Development in Nephrology

• Use of clinical trials with **master protocols**, a modern approach to expedite drug development

• Master protocols (basket trials, umbrella trials and platform trials) are novel designs that have the potential to explore multiple hypotheses through concurrent sub-studies

• The oncology field has pioneered these novel study designs

• A Basket Trial involves a single investigational drug (or drug combination) that is studied across multiple disease populations (can be defined by stage, histology, number of prior therapies, genetic or other biomarkers, demographic characteristics, etc.)
Advantages of Basket Trials

• Ideal for studying multiple diseases that have a common target

• Leverages the efficiency of one master protocol and one study system infrastructure

• The sub-studies within basket trials are usually designed as single-arm activity-estimating, in nephrology likely looking at proteinuria reduction or eGFR

• Each sub-study may have different objectives and endpoints

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Master Protocols: Efficient Clinical Trial Design Strategies to Expedite Development of Oncology Drugs and Biologics Guidance for Industry, FDA Draft Guidance 2018
Atrasentan in a Basket Trial for Treatment of Glomerular Diseases
ET\textsubscript{A} receptor activation drives proteinuria, mesangial cell activation & kidney inflammation & fibrosis, all hallmarks of glomerular diseases

Intense kidney ET-1 & ET\textsubscript{A} receptor immunostaining in IgAN patients with significant proteinuria

ET\textsubscript{A} system activation appears to be a key molecular determinant of the clinical course in glomerular diseases

Elevated kidney ET-1 expression strongly & prospectively predicted progression of IgAN, 12 months following kidney biopsy

Blockade of the ET\textsubscript{A} receptor through atrasentan represents a promising approach to treat glomerular disease
Topline Results from SONAR

* * *

Atrasentan and renal events in patients with type 2 diabetes and chronic kidney disease: a double-blind, randomized, placebo-controlled trial

- **3600** high-risk patients with DKD randomized
- **2** Years median treatment duration
- **35%** decreased risk of ESKD or doubling of serum creatinine in responders* (28% in all randomized)
- **27%** risk reduction of End Stage Kidney Disease
- **39%** risk reduction of doubling of serum creatinine

![Graph showing the hazard ratio and proportion of patients at risk over time for Placebo and Atrasentan.](image)

Hazard ratio 0.65 (95% CI 0.49-0.88); *P* = 0.0047

<table>
<thead>
<tr>
<th>Time (m)</th>
<th>Number at risk (Placebo)</th>
<th>Number at risk (Atrasentan)</th>
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</table>

*Heerspink et al, Diabetes Obes Metab. 2018;20:1369–1376, Heerspink et al, SONAR Trial, Lancet 2019*
SONAR Outcomes Supportive of Success in Other Proteinuric Glomerular Diseases

Proteinuria reduction is recognized as an important surrogate endpoint in glomerular diseases

*In Responders (patients who achieved >30% reduction in proteinuria)

![Graph showing Proteinuria (UACR) and Kidney Function (eGFR Slope)]

Heerspink et al, Diabetes Obes Metab. 2018;20:1369–1376, Heerspink et al, SONAR Trial, Lancet 2019
AFFINITY: A Basket Trial of Glomerular Diseases
AFFINITY Study Overview

Study Objective

• The Phase 2 AFFINITY study assesses the efficacy and safety of atrasentan in patients with proteinuric glomerular diseases at risk of progressive kidney function loss

Study Design

• Approximately 80 patients (~20 per cohort) in the United States, Australia, South Korea, Italy, Spain and the United Kingdom with the following proteinuric glomerular diseases will receive 0.75 mg atrasentan for 52 weeks:
  • Cohort 1: IgAN
  • Cohort 2: FSGS
  • Cohort 3: Alport Syndrome
  • Cohort 4: DKD (on stable dose of SGLT2i)
• Patients must be on a maximally tolerated and stable dose of a RASi
• Where allowed by local regulations, options for remote study visits using telemedicine and home health may be offered
• Provides a flexible solution for patients and clinicians in the era of COVID-19 and reduces the burden to patients for trial participation
Methods

Study Objective

- AFFINITY is a global, phase 2, open label basket study to assess the efficacy and safety of atrasentan in patients with proteinuric glomerular diseases at risk of progressive kidney function loss

**Screening**

- IgA Nephropathy
  - (UPCR 0.5 < 1.0 g/g)
  - N=20

- FSGS
  - (UPCR > 1.5 g/g)
  - N=20

- Alport Syndrome
  - (UPCR > 0.5 g/g)
  - N=20

- Diabetic Kidney Disease (+ SGLT2i)
  - (UACR > 0.5 g/g)
  - N=20

**Atrasentan 0.75 mg daily for 52 weeks**

**Follow Up 56 weeks**

**Proteinuria Endpoint**

**Week 12**

**Exploratory eGFR Endpoint**

**Week 52**

*All cohorts eGFR ≥30 ml/min/1.73m², except for DKD ≥ 45 ml/min/1.73m²*
The primary endpoint is change in proteinuria from baseline at Week 12:

- Cohort 1-3: UPCR from a 24-hr urine collection
- Cohort 4: UACR from a First Morning Void

Additional outcome measures include:

- Evaluate atrasentan plasma concentration over time
- Cohort 1: Percent of subjects achieving proteinuria reduction to less than 0.3 g/day
- Cohort 2: Percent of subjects achieving UPCR < 1.5 g/g and > 40% reduction from baseline
- Cohort 1-4: Change from baseline to Week 52
- Cohort 1-4: Change from baseline in estimated glomerular filtration rate (eGFR) at Week 56
Inclusion Criteria, IgA Nephropathy

Subjects must meet ALL inclusion criteria to be enrolled

Types of Subjects and Disease Characteristics

Cohort 1- IgAN

Biopsy-proven IgAN that, in the opinion of the Investigator, is not due to secondary causes

- Biopsy could have occurred at any point in time prior to study
- A diagnostic report must be available for review by the Sponsor or designee

Receiving a maximally tolerated and optimized dose of a RAS inhibitor that has been stable for at least 12 weeks prior to screening

UPCR ≥ 0.5 and < 1.0 g/g (≥ 500 mg/g and < 1000 mg/g) based on a central laboratory assessment of first morning void urine collected at screening

eGFR ≥ 30 mL/min/1.73 m²
Inclusion Criteria, FSGS

Subjects must meet ALL inclusion criteria to be enrolled

Types of Subjects and Disease Characteristics

**Cohort 2 - FSGS**

Biopsy-confirmed FSGS or documentation of a genetic mutation in a podocyte protein associated with FSGS

UPCR > 1.5 g/g (>1500 mg/g) based on a central laboratory assessment of first morning void urine collected at screening

eGFR ≥ 30 mL/min/1.73 m²

Receiving a maximally tolerated and optimized dose of a RAS inhibitor that has been stable for at least 12 weeks prior to screening

If receiving systemic corticosteroids or calcineurin inhibitors, dose (level) must be stable for 12 weeks prior to start of study drug and anticipated to remain on a stable dose at least through week 12

Body Mass Index (BMI) ≤ 40 kg/m²
Inclusion Criteria, Alport Syndrome

Subjects must meet ALL inclusion criteria to be enrolled

Types of Subjects and Disease Characteristics

**Cohort 3 - Alport syndrome**

Diagnosis of Alport syndrome by genetic testing (documented mutation in a gene associated with Alport syndrome, including COL4A3, COL4A4, or X-linked COL4A5 in the subject or a family member) OR patients with a new mutation that, in the opinion of the Investigator, has significant supporting evidence of Alport syndrome (biopsy, familial genetics, family history & familial biopsy, microscopic hematuria, hearing loss pattern, fleck retinopathy)

UPCR > 0.5 g/g (>500 mg/g) based on a central laboratory assessment of first morning void urine collected at screening

Receiving a maximally tolerated and optimized dose of a RAS inhibitor that has been stable for at least 12 weeks prior to screening

eGFR ≥ 30 mL/min/1.73 m²
Inclusion Criteria, DKD

Subjects must meet ALL inclusion criteria to be enrolled

Types of Subjects and Disease Characteristics

Cohort 4- DKD

Clinical diagnosis of type 2 diabetes mellitus (T2DM) as per guidelines

Diagnosis of DKD based on the opinion of the investigator, including the presence of the following criteria:

a. UACR ≥ 0.5 g/g (500 mg/g) based on a central laboratory assessment of first morning void urine collected at screening
b. eGFR ≥ 45 mL/min/1.73 m²

Receiving a maximally tolerated and optimized dose of a RAS inhibitor that has been stable for at least 12 weeks prior to the screening visit and stable dose of SGLT2 inhibitor for at least 12 weeks prior to screening
Main Exclusion Criteria

Subjects must meet NONE of the following exclusion criteria to be enrolled

Concurrent diagnosis of another cause of chronic kidney disease including diabetic kidney disease or another primary glomerulopathy

Clinical suspicion of rapidly progressive glomerulonephritis (RPGN) based on KDIGO guidelines or clinical suspicion of IgA vasculitis (Henoch-Schonlein Purpura)

Known history of congestive heart failure, diastolic dysfunction, or prior hospital admissions for conditions relating to fluid overload such as pulmonary edema, uncontrolled peripheral edema, pleural effusion, or ascites.

Confirmed blood pressure >150 mmHg systolic or >95 mmHg diastolic based on a mean of 3 measurements obtained at screening.

With the exception of DKD (Cohort 4), use of an SGLT2 inhibitor within the past 30 days.

HbA1c > 9.5% in Cohort 4 (DKD), HbA1c > 7.0% in Cohorts 1-3.

Except for Cohort 2 (FSGS), use of systemic immunosuppressant medications including systemic steroids (prednisone or equivalent >10 mg/day for more than 2 weeks within 3 months prior to screening), mycophenolate, azathioprine, cyclosporine, tacrolimus, etc. for more than 2 weeks within the past 3 months prior to screening.
Chinook plans to present data from the IgAN patient cohort of the AFFINITY study in the first half of 2022!

Stay tuned!
Chinook Booth at the 59th ERA Congress

The 59th ERA Congress will take place on May 19-22, 2022, both virtual and live in Paris.

Visit Chinook at our exhibit booth!
Join Us in our Mission to Discover and Develop Precision Therapies for Patients with Kidney Diseases

Contact us if you would like to learn more about any of our trials, would like to refer a patient or are interested in becoming one of our investigators!

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