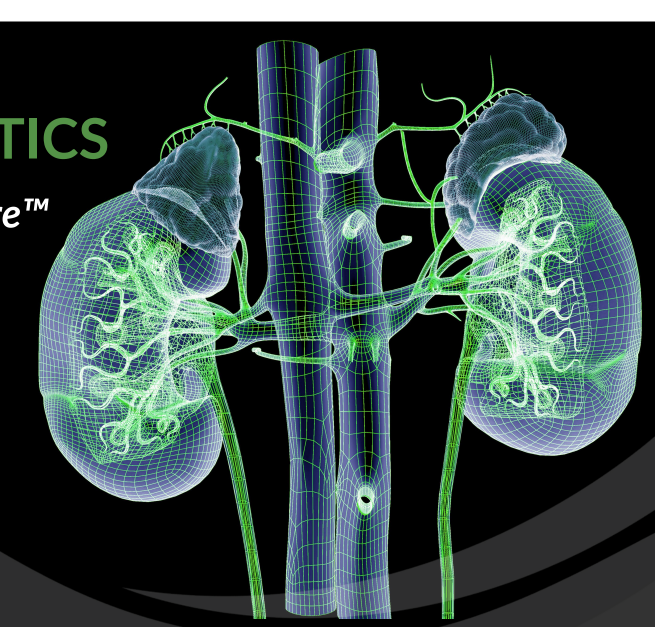


CHINOOK THERAPEUTICS

Changing the Course of Kidney Care™

March 2023



## **Note Regarding Forward-Looking Statements**



Certain of the statements made in this presentation are forward looking, including those relating to Chinook's business, future operations, advancement of its product candidates and product pipeline, clinical development of its product candidates, including expectations regarding timing of initiation and results of clinical trials and sufficiency of its cash resources. In some cases, you can identify these statements by forward-looking words such as "may," "will," "continue," "anticipate," "intend," "could," "project," "expect" or the negative or plural of these words or similar expressions. Forwardlooking statements are not guarantees of future performance and are subject to risks and uncertainties that could cause actual results and events to differ materially from those anticipated, including, but not limited to, our ability to develop and commercialize our product candidates, whether results of early clinical trials or preclinical studies will be indicative of the results of future trials, our ability to obtain and maintain regulatory approval of our product candidates, our ability to operate in a competitive industry and compete successfully against competitors that may be more advanced or have greater resources than we do, our ability to obtain and adequately protect intellectual property rights for our product candidates. Many of these risks are described in greater detail in our filings with the SEC. Any forward-looking statements in this presentation speak only as of the date hereof. Chinook assumes no obligation to update forward-looking statements whether as a result of new information, future events or otherwise, after the date of this presentation.



CHRONIC KIDNEY DISEASES ARE A

# SEVERE & GROWING

**WORLDWIDE PROBLEM** 

KIDNEY DISEASE AFFECTS

10%

OF GLOBAL POPULATION

800M people worldwide, including 37M in U.S.<sup>1, 2</sup>

STAGGERING COSTS TO HEALTHCARE SYSTEM

>\$130B

Annual U.S. healthcare costs driven by kidney diseases<sup>3</sup>

KIDNEY DIALYSIS



156

Average hospital and dialysis center visits a year/patient<sup>4</sup>



\$200K

Per patient annually<sup>5</sup>

KIDNEY TRANSPLANT



~23K

Transplants annually (U.S)<sup>6</sup>



\$400K

Per transplant<sup>7</sup>



4 YRS

Waitlist (U.S.)6



## THE TIME IS NOW FOR

# KIDNEY DISEASE DRUG DEVELOPMENT

#### **CLEAR DEVELOPMENT PATHS**

- | Increased understanding of underlying disease biology
- New and more validated drug targets
- FDA now recognizes surrogate markers, such as proteinuria and eGFR, as registration endpoints<sup>1</sup>

SHIFT AT THE FDA ALLOWS FOR MORE EFFICIENT STUDIES

5+ YRS



2-3 YRS

Traditional hard kidney outcomes trials require large numbers of patients and take many years to complete

Surrogate endpoint studies can be much smaller, with potential for accelerated approval in 6-9 months and full approval in 2-3 years

## VAST SHORTAGE OF NEW KIDNEY DRUGS

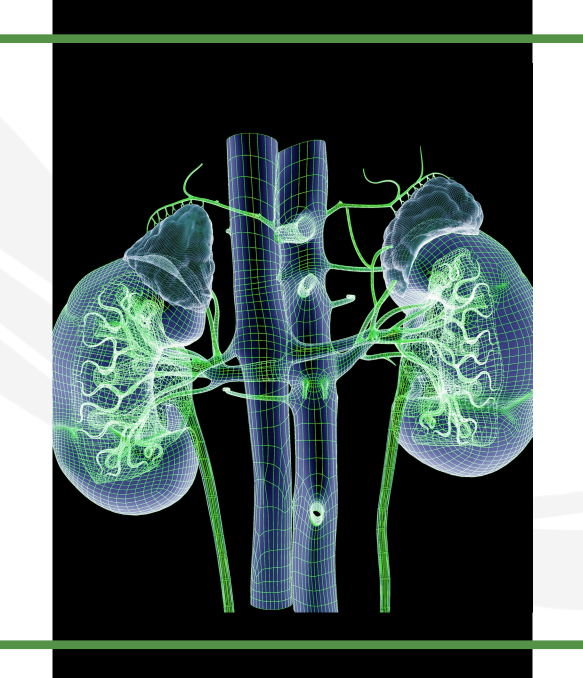
IN THE PAST 2 YEARS THERE HAVE BEEN

5 approved drugs for kidney diseases<sup>2</sup>

CURRENTLY, THERE ARE

40+ late-stage drugs in development<sup>3</sup>





# WE ARE CHANGING THE COURSE OF KIDNEY CARE™











## Advancing a Diversified Pipeline of Best-in-Class Programs



Program		Indication	Target Validation	Lead Optimization	IND-Enabling	Phase 1	Phase 2	Phase 3
_	ALIGN	IgA Nephropathy			Phase 3 ongoing with	ı topline data e	expected in H2	2023
Atrasentan	AFFINITY	Basket of glomerular diseases				Phase 2 or	ngoing	
BION-1301		IgA Nephropathy		Phase 1/2 ongoing	g with phase 3 planned	to start in mid	-2023	
СНК-336		Primary & Idiopathic Hyperoxalurias			Phase 1 HV study ong	going		
Research & Discovery Programs		Rare, severe chronic kidney diseases		Multiple programs				

Continuing to evaluate opportunities to add kidney disease programs to pipeline



## IgA NEPHROPATHY HAS A

## LARGE UNMET MEDICAL NEED

IgAN is the most common primary glomerular disease globally and requires chronic treatment

~150,000

Biopsy-confirmed IgAN patients in the U.S.<sup>1</sup>



~45% with >1 g/day<sup>1</sup>

~25% with 0.5 - 1 g/day1

Patients with uncontrolled proteinuria despite optimized standard of care RAS inhibition (ACEi or ARB)

~100,000 Patients remain at high risk for progression

High-risk patients living with IgAN will cycle through multiple therapies over their disease course, resulting in several million patient years of treatment, which will increase as time to ESKD is further delayed

~30-45% of IgAN patients progress to ESKD over 20-25 years<sup>3-6</sup>

**ACHIEVING** 

30% PROTEINURIA REDUCTION

**EQUATES TO** 

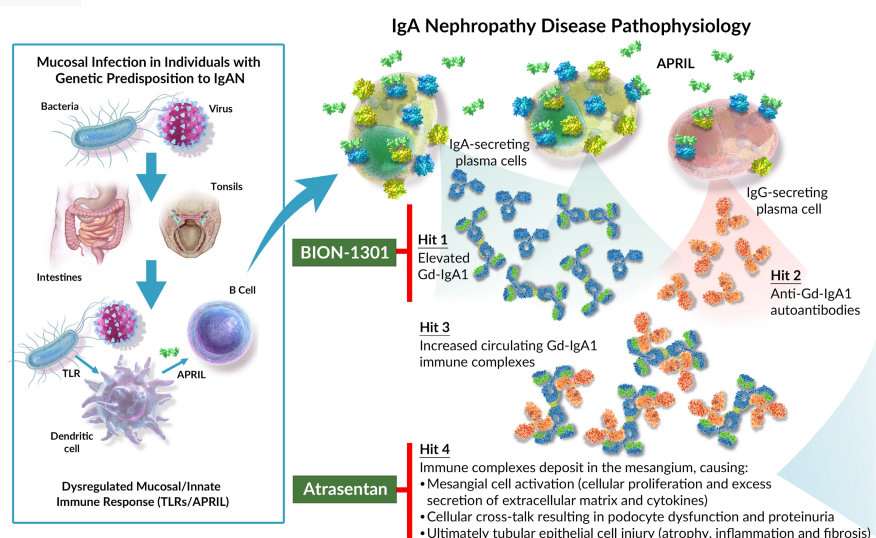
TIME TO ESKD<sup>2</sup>

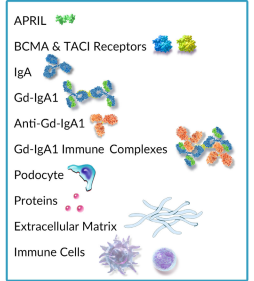
**Greater proteinuria reductions** are associated with greater clinical benefit

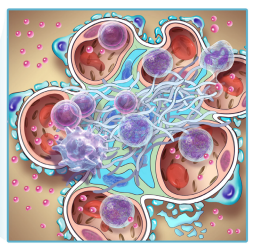


## Atrasentan & BION-1301: Two Complementary Programs for IgA Nephropathy

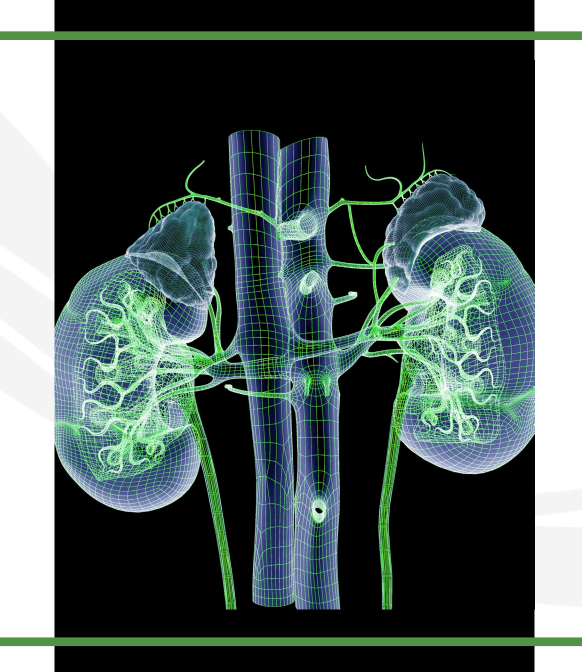
















**BION-1301** 

Anti-APRIL Monoclonal Antibody



CHK-336
LDHA Inhibitor



**R&D PROGRAMS** 

Precision Medicine Pipeline



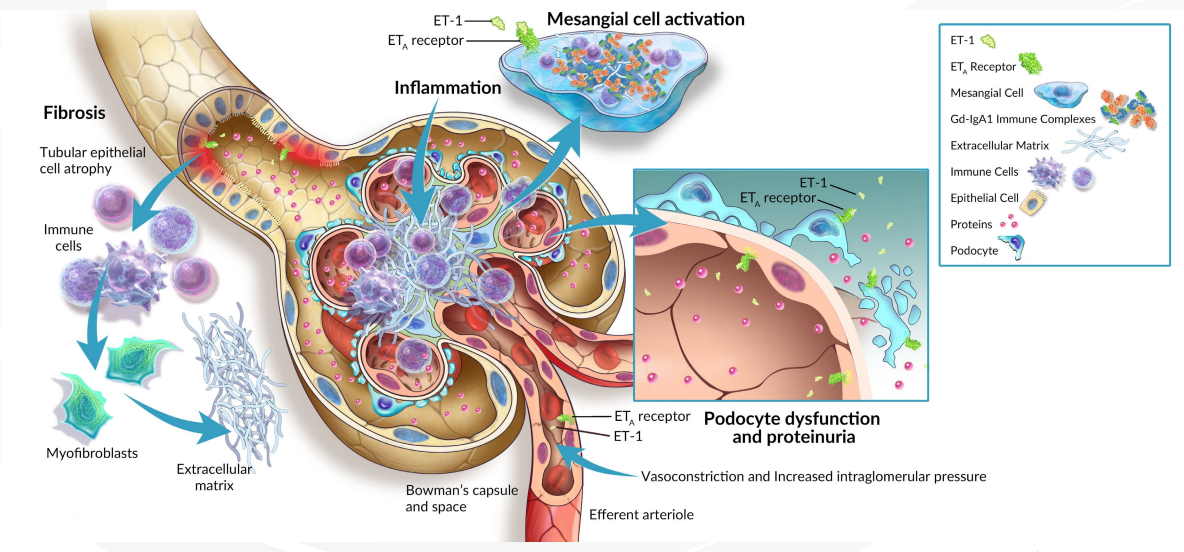
FINANCIALS & CATALYSTS

2023



## Atrasentan: A Potent and Selective ET<sub>A</sub> Antagonist ET<sub>A</sub> Receptor Activation Drives the Hallmarks of IgAN Progression Through Multiple Mechanisms







## Phase 2 AFFINITY™ Basket Study



### **Exploring Potential Across Proteinuric Glomerular Diseases**

#### TRIAL DESIGN:

- Open-label
- 12-week proteinuria primary endpoint
- 52-week treatment period
- 20 patients per cohort

### **COHORTS INCLUDE:**

- IgAN with proteinuria 0.5 <1 g/g
- Two FSGS cohorts (with dose escalation to 1.5 mg)
- Alport syndrome
- DKD combined with SGLT2 inhibitors

### Phase 2 AFFINITY™ IgAN Cohort Interim Results:

- >50% mean proteinuria reduction at 24 weeks
- Well-tolerated with no treatment-related severe AEs to date
- No weight gain or other evidence of significant fluid retention

### 

Median (Q1, Q3) baseline protein excretion: 1.2 (0.9, 1.5) g/day

20

20





19

## Phase 3 ALIGN™ Study in IgA Nephropathy



## Phase 3 Targeting IgAN Patients at High Risk for Disease Progression

- Biopsy-proven IgAN
- Patients on maximally-tolerated, optimized and stable dose of RASi, or RASi intolerant
- Proteinuria >1 g/day and eGFR >30 ml/min/1.73m<sup>2</sup>
- ~320 pts, 1:1 placebo randomization
- Global study at ~170 sites
- Primary endpoint: 24-week proteinuria (accelerated approval)
- Secondary endpoint: 2.5-year eGFR (full approval)

## Two-Pronged Approach to Evaluate Atrasentan in Combination with SGLT2i

## ALIGN™ SGLT2i + Atrasentan Combination Stratum Underway

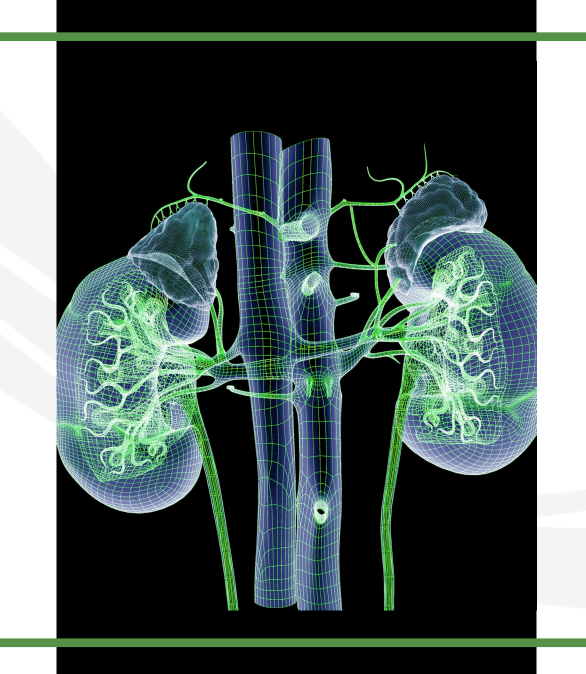
- Stratum of up to 64 patients in ALIGN™ on stable dose of SGLT2i
- Safety and efficacy (proteinuria + eGFR) exploratory analysis with no change to enrollment timelines or primary analysis population

## Planned Phase 2 Study of SGLT2i + Atrasentan Combination in IgAN

- Randomize patients 1:1 to atrasentan or placebo
- Primary endpoint: change in UPCR from baseline to week 12
- Data to support future use of atrasentan in combination with SGLT2is

2020	2021	2022	2023	2024	2025	2026	2027
J:							
ALIGN			Top line proteinuria data		eGFR endpoint		
			in H2 2023		da	ta	









Anti-APRIL Monoclonal Antibody



**CHK-336** 

**LDHA** Inhibitor



## **R&D PROGRAMS**

Precision Medicine Pipeline



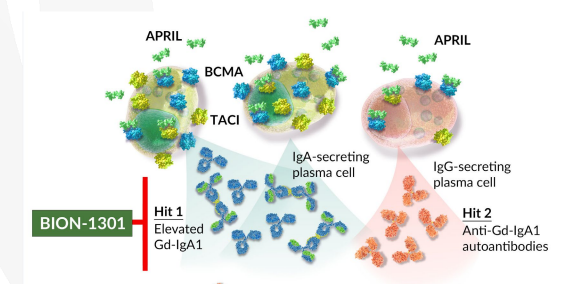
## FINANCIALS & CATALYSTS

2023



## **BION-1301: Potentially Disease-Modifying Anti-APRIL mAb** in IgAN





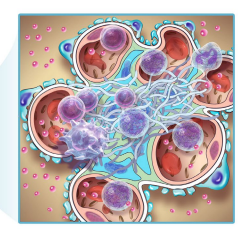
### Hit 3 Increased

circulating Gd-IgA1 immune complexes

#### Hit 4

Immune complexes deposit in the mesangium, causing:

- Mesangial cell activation (cellular proliferation and excess secretion of extracellular matrix and cytokines)
- Cellular cross-talk resulting in podocyte dysfunction and proteinuria
- Ultimately tubular epithelial cell injury (atrophy. inflammation and fibrosis)



### **APRIL**

TNF-family cytokine involved in B-cell signaling<sup>1</sup>

- **Drives IgA production** and survival of IgA-secreting plasma cells<sup>2</sup>
- Shown to increase Gd-IgA1 secretion<sup>3</sup>
- Higher APRIL levels in IgAN patients correlated with higher Gd-IgA1 and proteinuria and lower eGFR<sup>3</sup>
- APRIL gene variants confer increased risk of IgAN<sup>4</sup>

### **BION-1301**

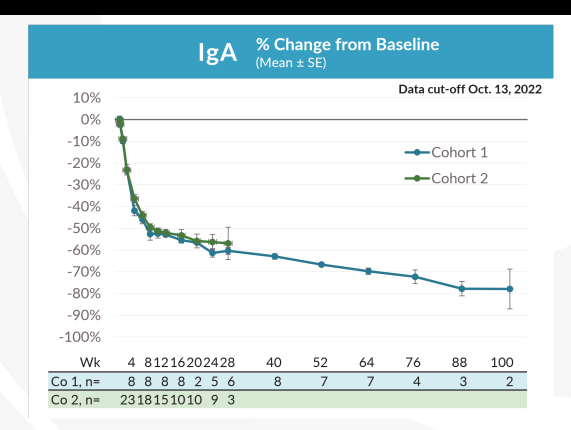
humanized IgG4 monoclonal antibody that blocks APRIL binding to its receptors

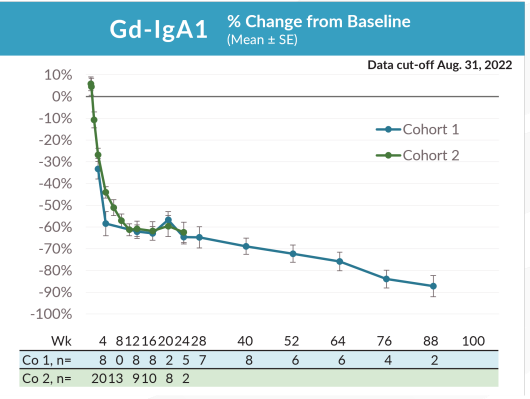
- Well-tolerated to date in:
  - NHPs<sup>5</sup>
  - Healthy volunteers<sup>7</sup>
  - Patients with multiple myeloma at doses up to 2700mg<sup>6</sup>
  - Patients with IgAN<sup>8</sup>
- Demonstrated disease-modifying potential in patients with IgAN<sup>8</sup>



## BION-1301 Results in Rapid and Durable Reductions in IgA and Gd-IgA1 in Patients with IgAN







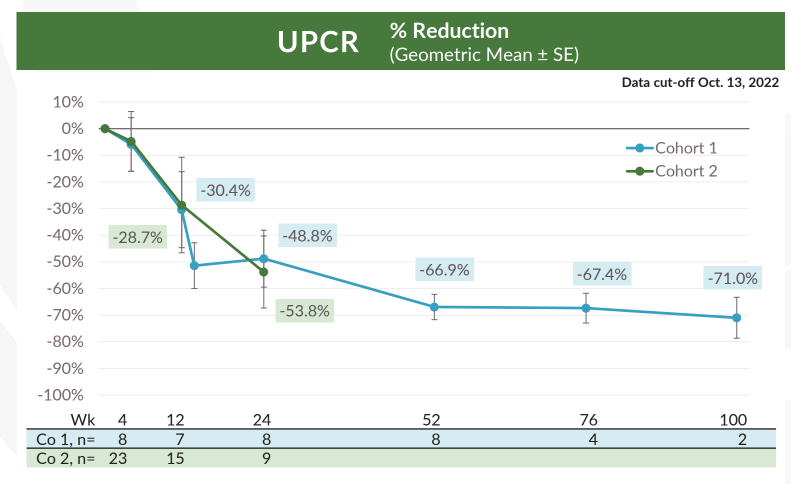
Mean Gd-IgA1 are not available at week 100

- Reductions in IgM, and to a lesser extent IgG, also observed
- BION-1301 generally well-tolerated in IgAN patients, with no reported deaths, SAEs, or AEs leading to discontinuation of study drug to date
- No ADAs observed to date



## BION-1301 Treatment Results in Sustained, Clinically Meaningful Proteinuria Reductions in Patients with IgAN





Median (min, max) baseline protein excretion: Cohort 1, 1.2 (0.7, 6.5) g/day; Cohort 2, 1.0 (0.6, 2.7) g/day

### COHORT 1 (IV $\rightarrow$ SC)

- Clinically meaningful reductions in UPCR were seen in patients with IgAN across a wide range of baseline proteinuria levels
- UPCR continued to decline through one year and was maintained through two years, providing evidence of sustained efficacy
- At Week 52, 7/8 evaluable patients demonstrated >50% reductions in UPCR

### COHORT 2 (de novo SC)

 Mean reduction in UPCR of >50% at 24 weeks in Cohort 2 with de novo SC administration is consistent with Cohort 1



## BION-1301 MOVING FORWARD

Plan to advance cohort 2 dose/schedule in pivotal trial, given strong clinical data

#### **STATUS**

Cohort 1 in IgAN

450 mg IV → 600 mg SC q2w
Enrollment of 10 Patients Completed

Cohort 2 in IgAN

600 mg SC q2w
Enrollment of 30 Patients Completed

## **NEXT STEPS**



Align with global health authorities (ongoing)



Conduct site and country feasibility (ongoing)



Initiate pivotal trial in mid-2023



## **Proposed BION-1301 Phase 3 Trial Design**

Initiate Pivotal Trial in Mid-2023, Given Strong Clinical Data and Disease-Modifying Potential



## Phase 3 Targeting IgAN Patients at Risk for Disease Progression

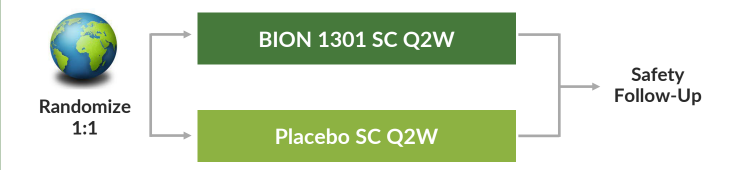
## **Key Inclusion Criteria**

- Biopsy-proven IgAN
- Patients on maximally-tolerated, optimized and stable dose of RASi (≥12w), or RASi intolerant
- Background optimized and stable dose (≥12w), of SGLT2i or ERA (if approved) allowed
- Proteinuria >1 g/day; eGFR >30 ml/min/1.73m²

## **Key Exclusion Criteria**

- Secondary IgAN, IgA vasculitis, other CKD, RPGN
- Recent immunosuppressant use, immune-deficient state, current severe infection, IgG < 6g/L</li>

Exploratory cohort eGFR 20 to < 30 mL/min/1.73m<sup>2</sup> (n~20)



### **Primary Endpoint**

UPCR @ 9 mos (40wks), n= 204

### **Additional Secondary Endpoints**

Composite 30% or 40% reduction in eGFR, eGFR < 15 mL, dialysis, kidney transplantation or all-cause mortality

Percent of subjects achieving a ≥ 25% reduction of UPCR to < 1.0 g/day at week 40

### **Key Secondary Endpoint**

eGFR (change from baseline) @ 2 yrs (104 wks), n=272

### **Safety Endpoints**

Type, incidence and severity of AEs and AESIs

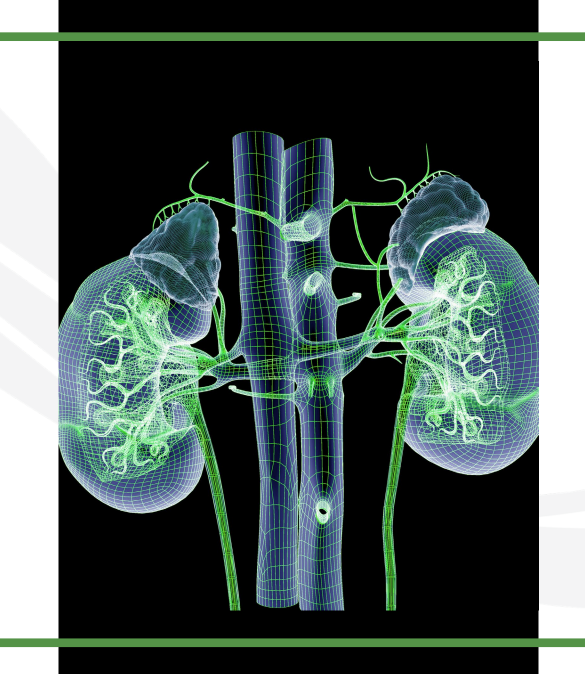
### **Exploratory Endpoints**

Characterize PK, exposure response, immunogenicity, QOL, MOA

Stratification Factors

Proteinuria (≥ 2 g/day vs. < 2 g/day), eGFR (≤ 45 v > 45 mL/min, Region (Asia v ROW)









**BION-1301** 

Anti-APRIL Monoclonal Antibody







FINANCIALS & CATALYSTS

2023



## Hyperoxalurias are Diseases Caused by Excess Oxalate

Hyperoxaluria is an Important Risk Factor for Kidney Stones

PRIMARY HYPEROXALURIAS (PH) 1-3 ARE ULTRA-RARE DISEASES

- Caused by genetic mutations resulting in overproduction of oxalate by the liver
- PH leads to recurrent kidney stones and can lead to kidney failure, if left untreated
- Median age of kidney failure for most severe form of disease
   (PH1) is 23 years<sup>1</sup>
- ~5,000 7,000 PH1 patients in the US and Europe<sup>2</sup>

#### IDIOPATHIC HYPEROXALURIA IS MORE COMMON

- May result from increased endogenous oxalate overproduction, particularly in association with metabolic diseases<sup>3</sup>
- Hyperoxaluria (generally defined as urinary excretion of >40 mg/d)
   is present in ~20–40% of frequent stone formers

## Decline in kidney function results in systemic oxalosis, affecting multiple organs<sup>4-7</sup>







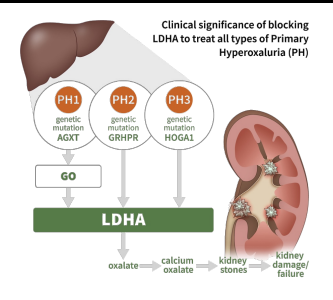




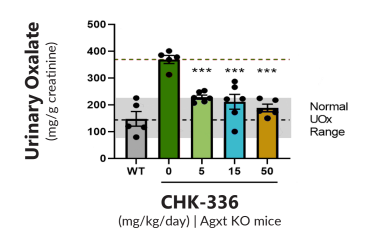
## CHK-336: Oral Small Molecule LDHA Inhibitor for PH

Liver-Targeted Tissue Distribution Profile Enables Potential to Treat All PH Types





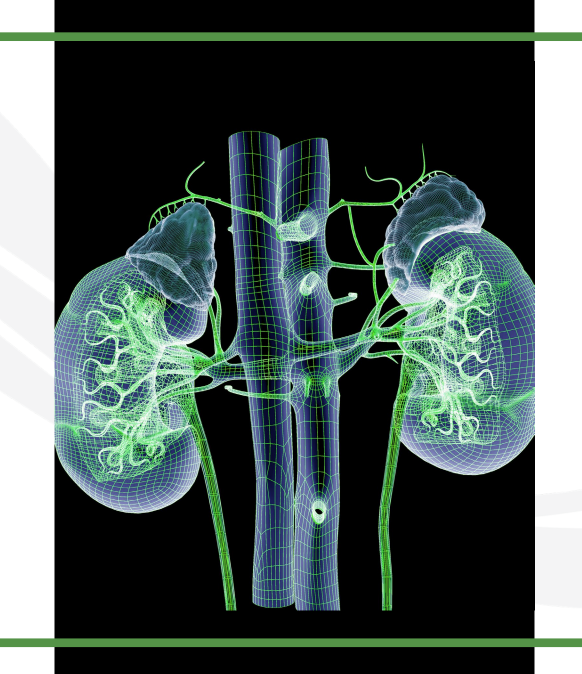
EFFICACY IN MOUSE PH1 MODEL



- Lactate dehydrogenase (LDHA) is the final step in production of oxalate from glyoxylate (GO) in the liver<sup>1</sup>
- Targeting LDHA may address all types of PH

- CHK-336 produces significant and dose-dependent urinary oxalate reductions in PH1 mouse models<sup>2</sup>
- Significantly reduces urinary oxalate excretion in a PH2 mouse model<sup>3</sup>
- Ongoing phase 1 SAD/MAD study evaluating safety, tolerability and PK of CHK-336 in healthy volunteers
- Anticipate reporting phase 1 healthy volunteer data in H1 2023
- Advancing towards initiation of phase 2 POC trials for patients with primary and idiopathic hyperoxalurias









**BION-1301** 

Anti-APRIL Monoclonal Antibody









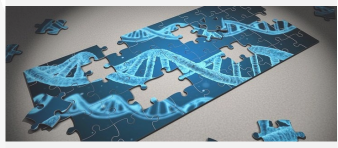
## Precision Medicine Approach to Research & Discovery

Focused on Rare, Severe CKDs with Defined Genetic and Molecular Drivers



### TARGET SELECTION & VALIDATION

### Systems Biology



Molecular Classification of CKD

- Target ID
- Target Validation
- Patient Stratification







### Translational Models



Modeling Human Disease

- Disease Mechanisms
- Target Validation
- Deep Biological Insights

Detailed insights into molecular pathogenesis of stratified CKDs

### TARGET EXECUTION

### **Development Candidates**



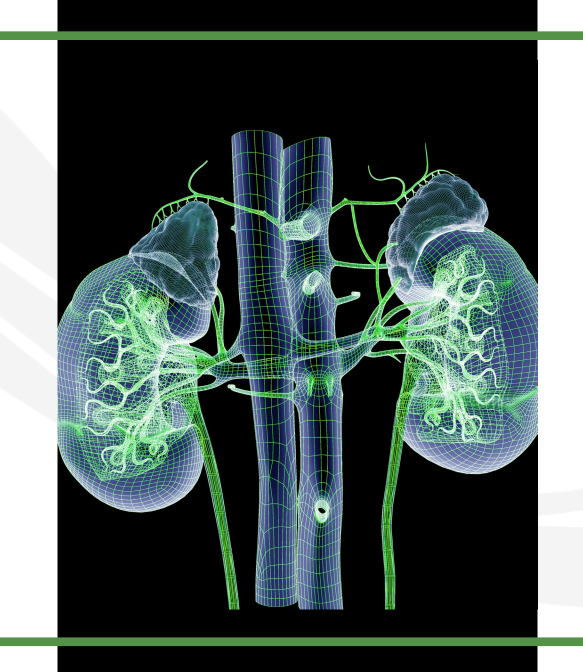


- First-in-Class or Best-in-Class
- Expert & Focused Chemistry,
   Biology, Pharmacology, DMPK,
   BD

Novel & differentiated molecules











**BION-1301** 

Anti-APRIL Monoclonal Antibody









# FINANCIAL STRENGTH NASDAQ: KDNY

STRONG BALANCE SHEET

\$385.3M

in cash, cash equivalents and marketable securities as of December 31, 2022

COMMON STOCK OUTSTANDING



~70.5 M

shares as of February 17, 2023\*



~75.3 M

fully diluted shares as of February 24, 2023\*\*

CASH GUIDANCE

Operating capital into 2025 based on current business plan

SANRENO JOINT VENTURE IN CHINA

- 50:50 JV for atrasentan and BION-1301
- Provides local execution and enhanced access to large IgAN patient populations in Asia
- Financial upside through equity ownership, milestones and royalties
- Reciprocal rights of first negotiation for future developed or in-licensed products



## Catalysts



	Program	Indication	Catalyst	H1 2023	H2 2023
BI		IgA Nephropathy	Initiate phase 2 trial in combination with SGLT2i in IgAN		
	Atrasentan		Report topline proteinuria data from ALIGN		
		Glomerular Diseases	Present additional data from other AFFINITY cohorts		
	BION-1301	IgA Nephropathy	Present additional phase 1/2 data from Cohorts 1 and 2 in IgAN		
	BION-1301-		Initiate phase 3 trial in IgAN in mid-2023		
	CHK-336	Hyperoxalurias	Report phase 1 healthy volunteer data and advance towards initiation of phase 2 POC trials in primary and idiopathic hyperoxalurias		





