

bridgebio

hope through
rigorous science

Corporate presentation

March 2021



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BridgeBio Pharma: Hope through rigorous science

Our mission: To **discover**, **create**, **test** and **deliver** transformative medicines to treat patients who suffer from genetic diseases and cancers with clear genetic drivers



Our 2025 vision – A leading player in genetic medicine



Multiple best-in-class or first-in-class products in blockbuster markets, with a total of 4+ NDAs on file

Patient-centric global commercial infrastructure

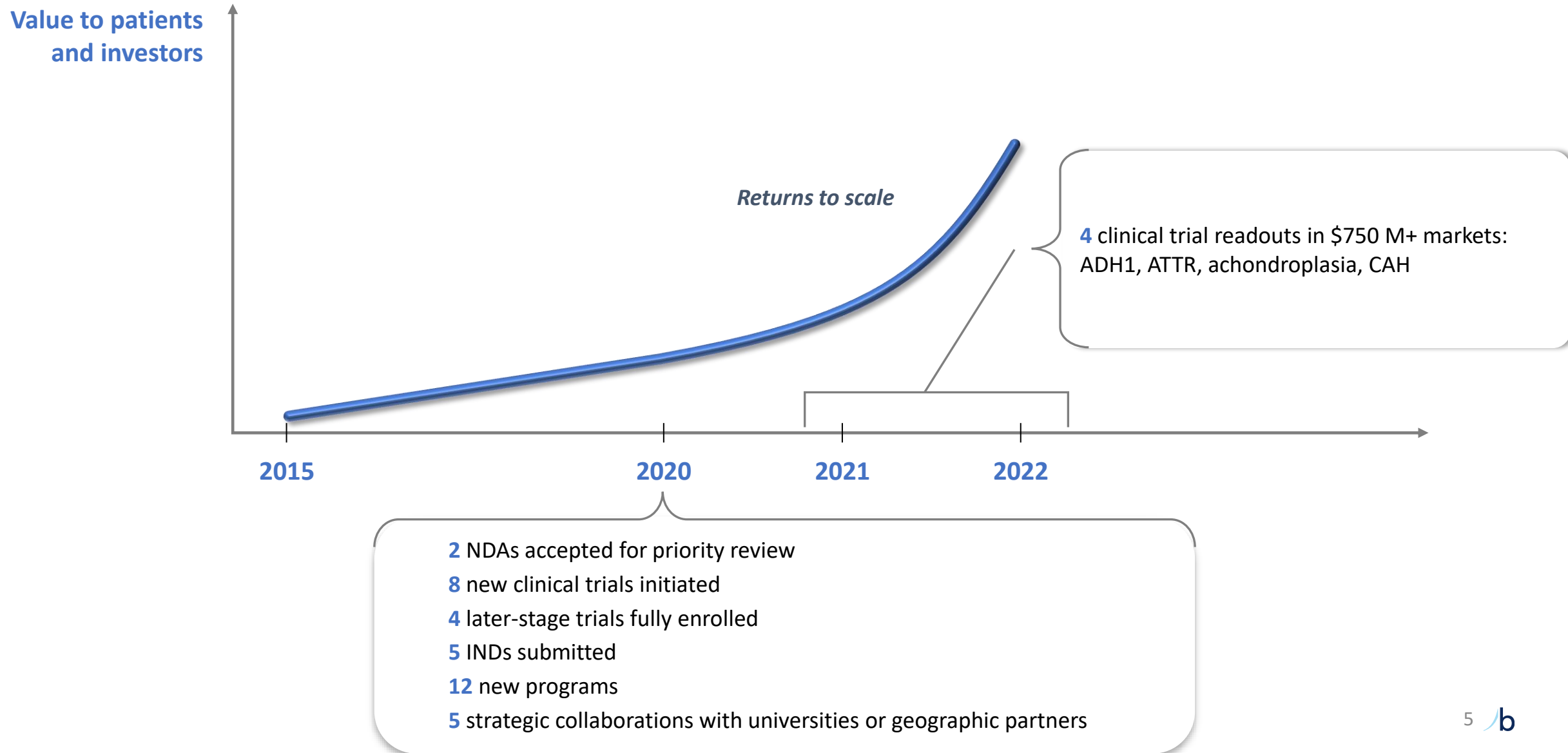
World-class drug discovery and development platform

Broad network of >40 university partnerships

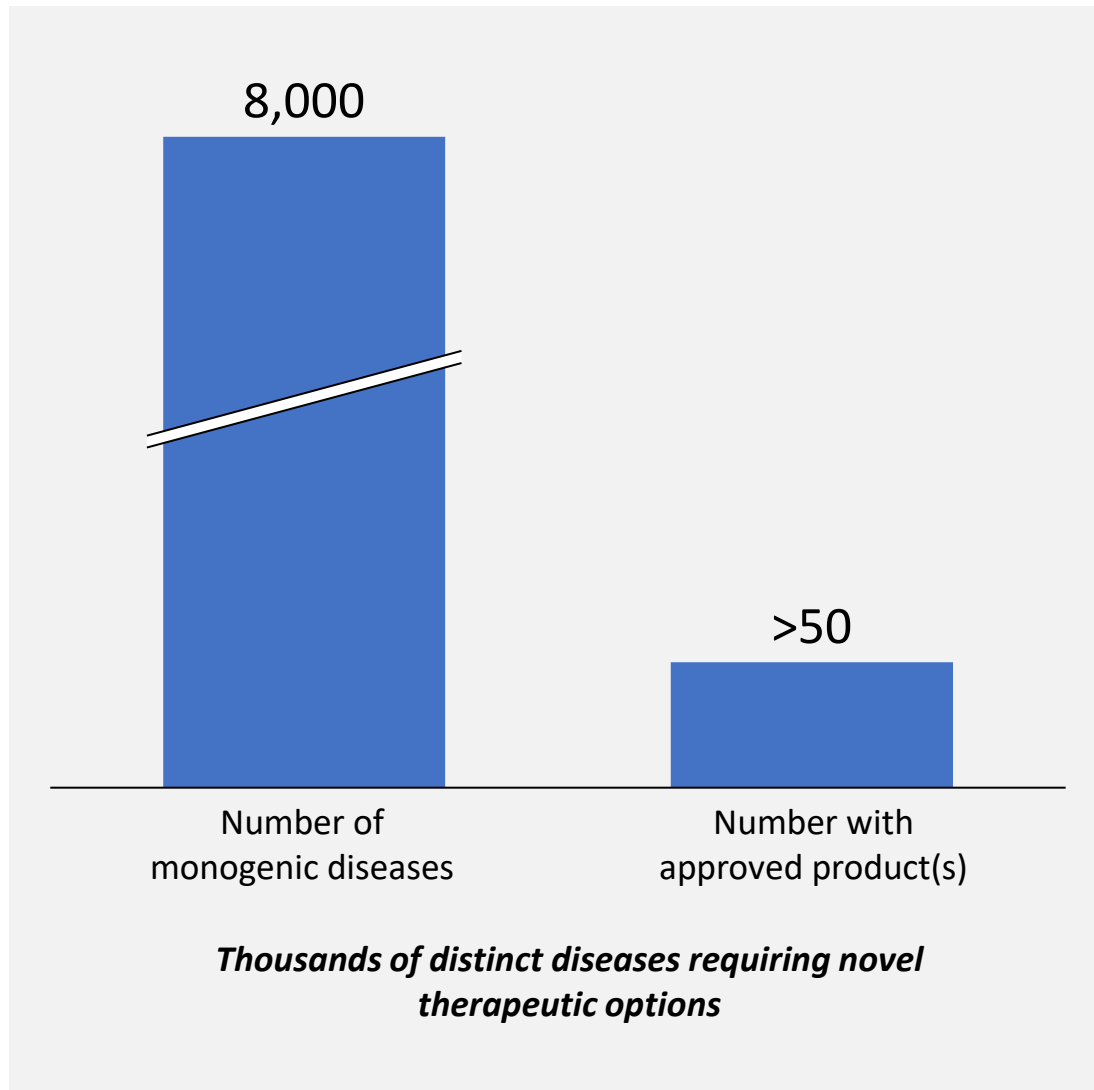
Multiple therapeutic modalities, many diseases

Deep pipeline of 30+ R&D programs

Context #1: 2021 is a critical year for BridgeBio



Context #2: The opportunity to help patients remains large



\$1B+
opportunities in
the pipeline

- 1) **Acoramidis** for ATTR CM and PN
- 2) **Low-dose infigratinib** for achondroplasia
- 3) **AAV5 gene therapy** for congenital adrenal hyperplasia
- 4) **High-dose infigratinib** for adjuvant urothelial carcinoma
- 5) **Pan-mutant KRAS inhibitor** for KRAS+ cancer
- 6) **SHP2 inhibitor** for RAS and kinase mutant cancer
- 7) **GPX4 inhibitor** for multiple tumor types
- 8) **GO1 inhibitor** for frequent kidney stone formers

Context #3: Still Day 1 for innovation within genetic medicine

Macromolecules

DNA

- gnomAD
- ENCODE3

RNA

- GTEx
- Single cell sequencing advances

Protein

- CryoEM
- DeepMind

Molecular Systems

- Mass spectrometry + metabolomics give us 1st snap of purine bio-synthesis

Clinical Diagnosis

- Whole genome sequencing of rare disease patients in UK Biobank
- Expanded sequencing led to novel causal variants in 28 genetic disorders

New Therapeutic Modalities

- Antisense oligonucleotides coming of age
- Gene therapy continues maturing

16 FDA approvals for drugs targeting rare genetic diseases or genetically defined cancers in 2020

Product platform: Our drug engineering platform leverages and efficiently translates innovation to therapies that matter

Discover

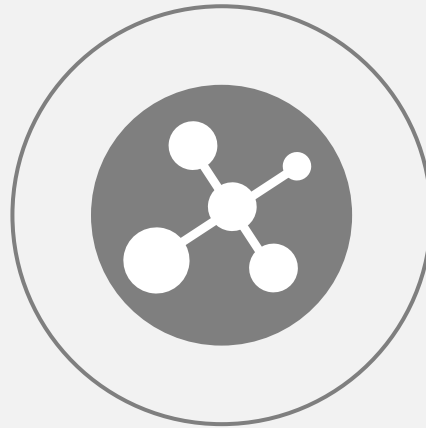
Novel genetic
disease targets



Computational genomics,
systemic disease mapping,
broad network of academic
partnerships

Create

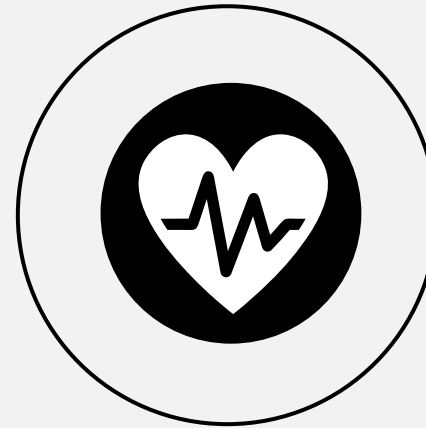
Medicines with industry-
leading research capabilities



Molecular dynamics assisted
chemistry, gene therapy,
therapeutic proteins, antisense
oligos

Test

Our drugs through global
development footprint



19 ongoing trials across >450
sites and 26 countries, central
operations toolkit and analytics

Deliver

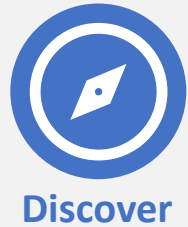
Our products to patients through
commercial infrastructure



Global infrastructure,
diagnostics, patient support,
disease state awareness

Product platform: BridgeBio is a people and a process

Scientific insight and judgment from industry leaders with a proven track record



Charles Homcy, MD
Founder and Chairman of
Pharmaceuticals



Frank McCormick, PhD
Founder and Chairman of
Oncology



Richard Scheller, PhD
Chairman of R&D



Len Post, PhD
Advisor



Phil Reilly, MD, JD
Advisor



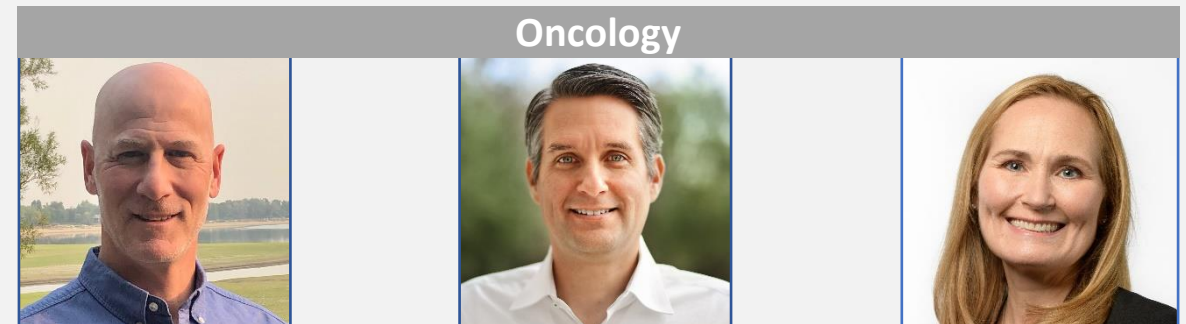
Experienced team of R&D operators responsible for 100+ INDs and 20+ approved products



Uma Sinha, PhD
Chief Scientific Officer



Robert Zamboni, PhD
Chemistry



Eli Wallace, PhD
Chief Scientific Officer,
Oncology



Pedro Beltran, PhD
SVP, Oncology





























Susan Moran, MD
Chief Medical Officer,
QED Therapeutics



Our pipeline spans multiple therapeutic areas with numerous upside opportunities

 Small molecule	 Topical small molecule	 Biologics	 Antisense oligo	 Gene therapy
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Portfolio segment	Program	Drug mechanism	Diseases	Patient pop. (US+EU)	Modality	Preclinical		Clinical		
						Discovery	IND-enabling	Phase1	Phase 2	Phase 3
Mendelian 	Acoramidis	TTR stabilizer	ATTR-CM	>400K						
	NULIBRY (fosdenopterin)	cPMP replacement	MoCD type A	100						Approved
	Infigratinib	Low-dose FGFR1-3i	Achondroplasia	55K						
	Encaleret	CaSR antagonist	ADH1 / HP	12K ¹ / 200K						
	BBP-418	Glycosylation substrate	LGMD2i	7K						
	BBP-711	GO1 inhibitor	PH1 / FSF	5K / 1.5M						
	BBP-671	PanK activator	PKAN / OA	7K						
	BBP-472	PI3Kβi	PTEN autism	120K						
	4 undisclosed small molecule programs			>500K						
	4 undisclosed antisense oligonucleotide programs			>300K						
Genetic Dermatology 	Patidegib²	Topical SMOi	Gorlin / BCC	120K						
	BBP-589	Recombinant COL7	RDEB	1.5K						
	BBP-681	Topical PI3Kai	VM / LM	117K						
	BBP-561	Topical KLK 5/7i	Netherton	11K						
Targeted Oncology 	Infigratinib	FGFR1-3i	3 FGFR+ tumor programs	37K						NDA filed
	BBP-398	SHP2i	Multiple tumors	>500K						
	BBP-454	Pan-mutant KRASi	3 KRAS+ tumors programs	>500K						
	BBP-954	GPX4i	Multiple tumors	>500K						
Gene Therapy 	BBP-631	21-OH gene therapy	CAH	>75K						
	BBP-812	ASPA gene therapy	Canavan	1K						
	BBP-815	TMC1 gene therapy	Genetic hearing loss	10K						
	4 undisclosed AAV gene therapy programs			150K						

¹ US carriers; ² We are party to an option agreement pursuant to which LEO Pharma A/S has been granted an exclusive, irrevocable option to acquire PellePharm, including the BBP-009 program. If the option is exercised by LEO Pharma A/S, we will no longer have rights to develop and commercialize BBP-009.

Product pipeline: Layers of de-risking and upside

Future pipeline catalysts and long-term growth

Targeted oncology
(FGFR3 in UC,
SHP2, KRAS)

Common mendelian
(LGMD2i, RDEB,
PKAN, VM)

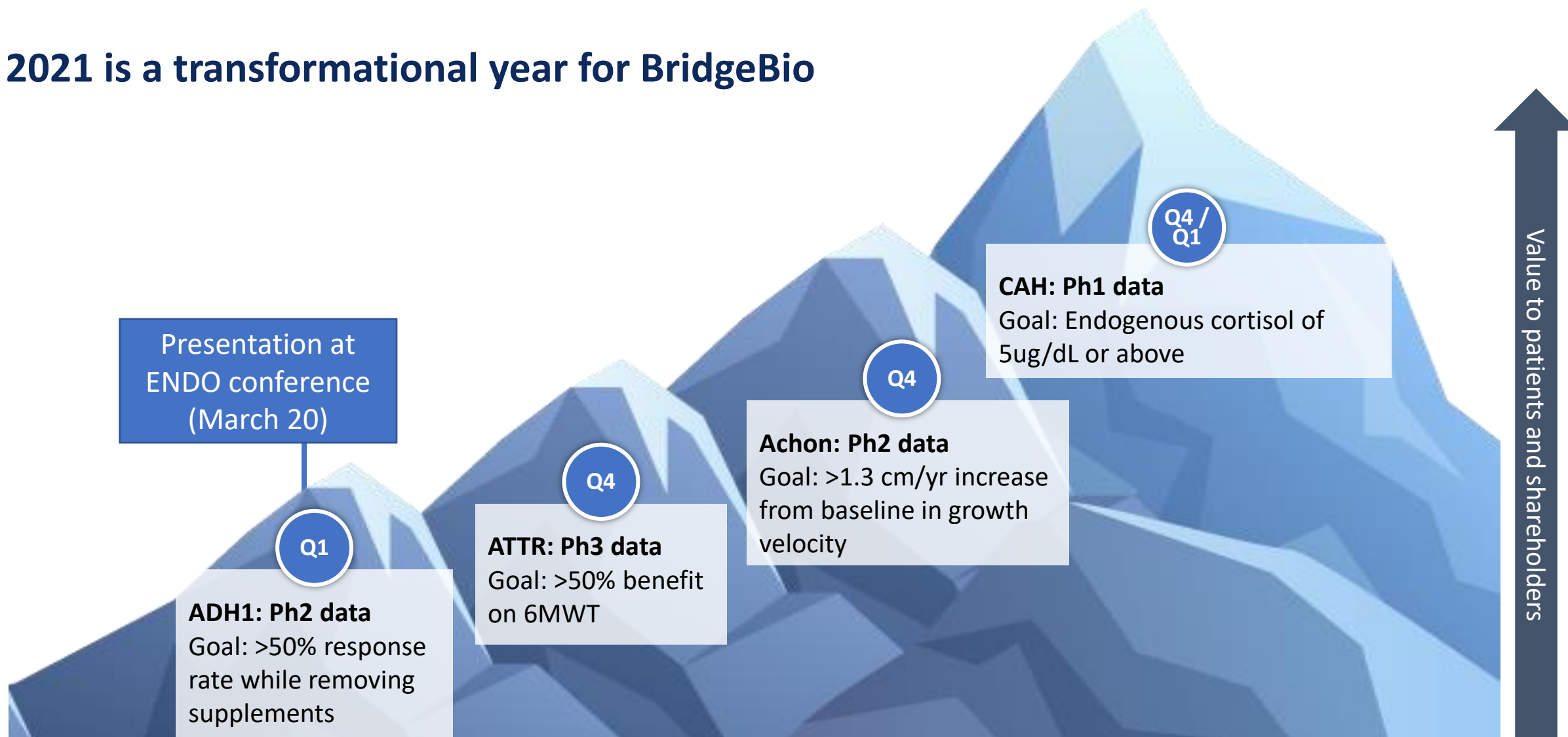
Validation of asset picking and execution

Near-term major catalysts from 4 core value drivers
(ATTR, ADH1, achondroplasia, CAH)

Proving ground and revenue

2 FDA approvals in 2021
(MoCD Type A, 2L+ CCA)

2021 is a transformational year for BridgeBio



Growth potential this year:

- Positive pivotal data in a multi-billion market
- Positive POC data in multiple blockbuster indications
- The right modality for the market and patients



Encaleret for autosomal dominant hypocalcemia type 1 (ADH1)

ADH1 overview



Prevalence

12-13K US variant carriers



Genetic driver

Calcium sensing receptor (CaSR) hyperactivation



Pathophysiology

Increased urinary calcium, decreased serum calcium and parathyroid hormone secretion

Features of a potential best-in-class medicine for ADH1



Direct targeting of CaSR

Normalization of all downstream effects of CaSR hyperactivity



Potential to address most **common symptoms**

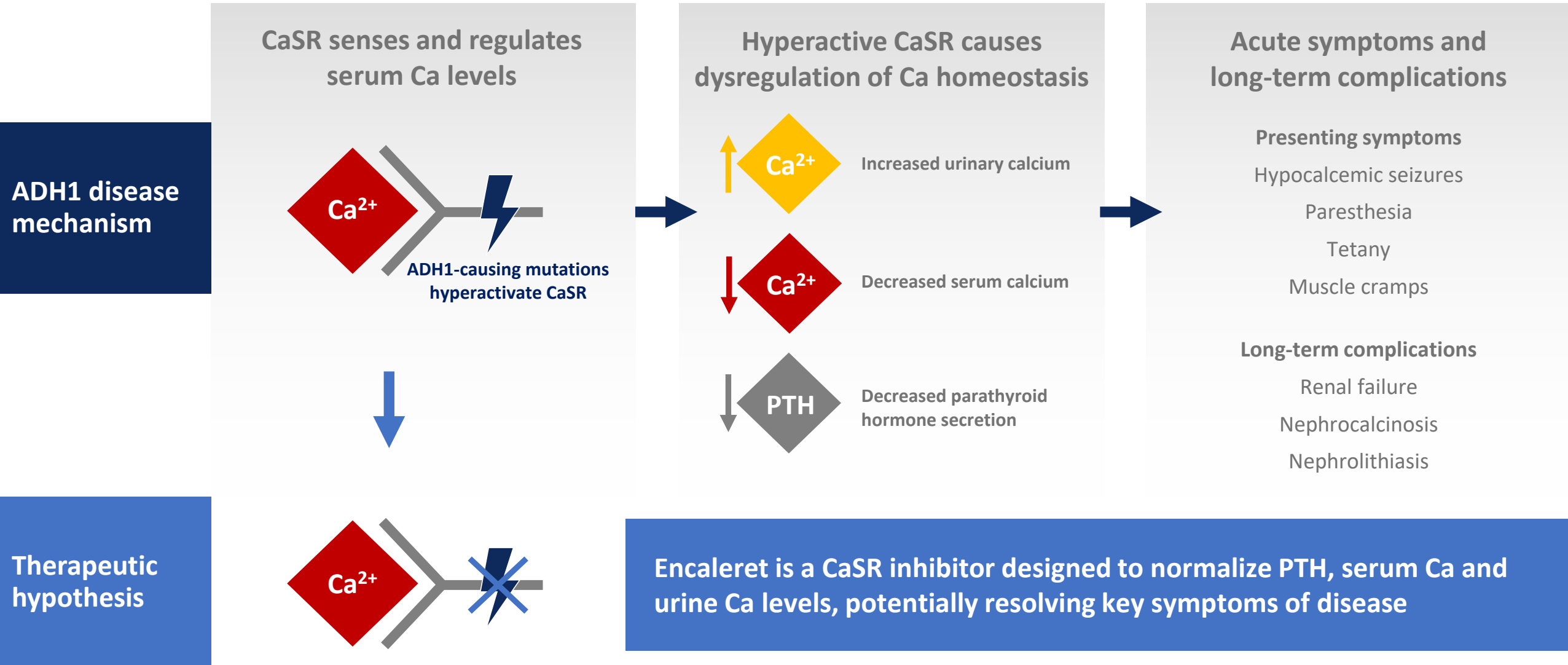
arising from altered calcium and parathyroid hormone dysregulation



Oral dosing, the first targeted therapy for ADH1 in a convenient form for patients and families

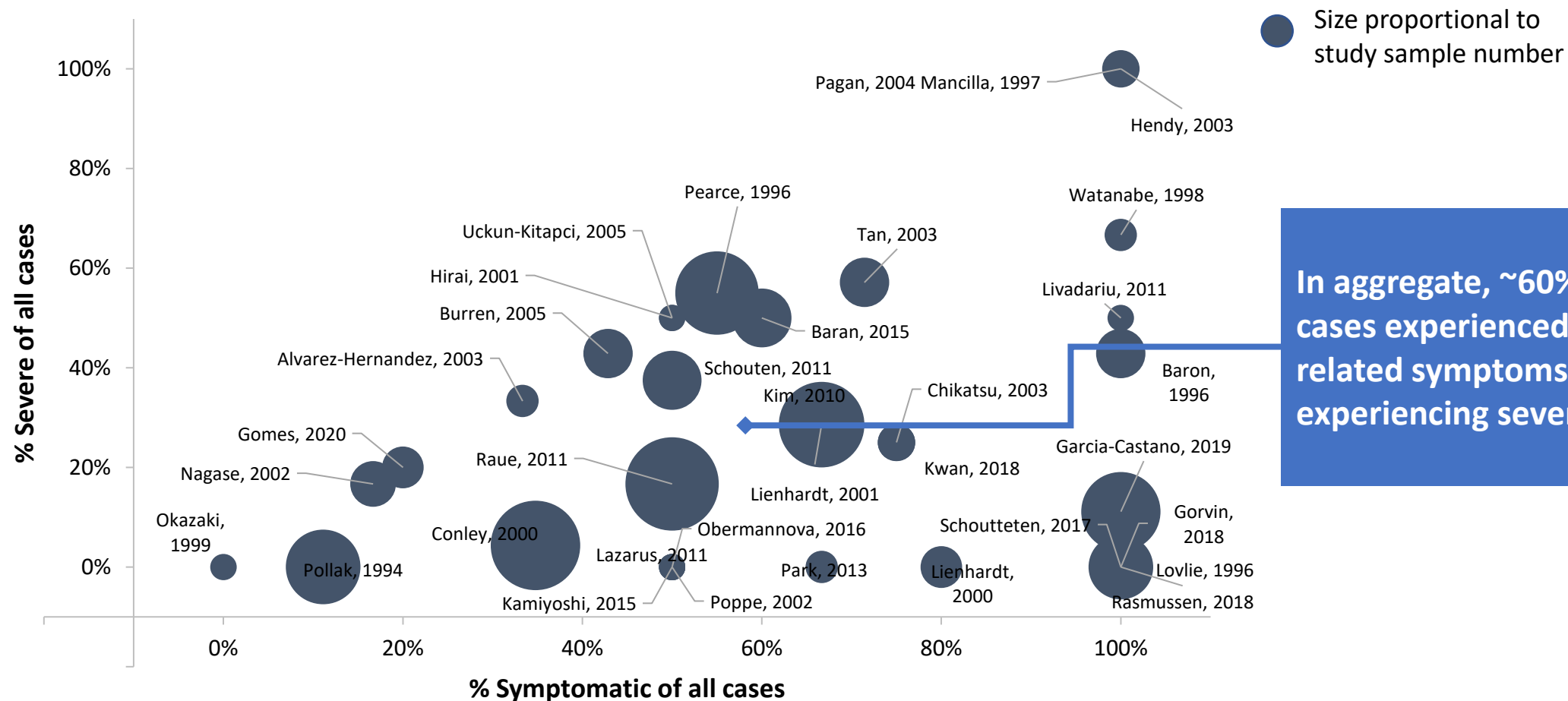
Alexis and Jackson
ADH1 patients

Encaleret is designed to treat ADH1 at its source



Majority of ADH1 patients are symptomatic including one third with severe symptoms

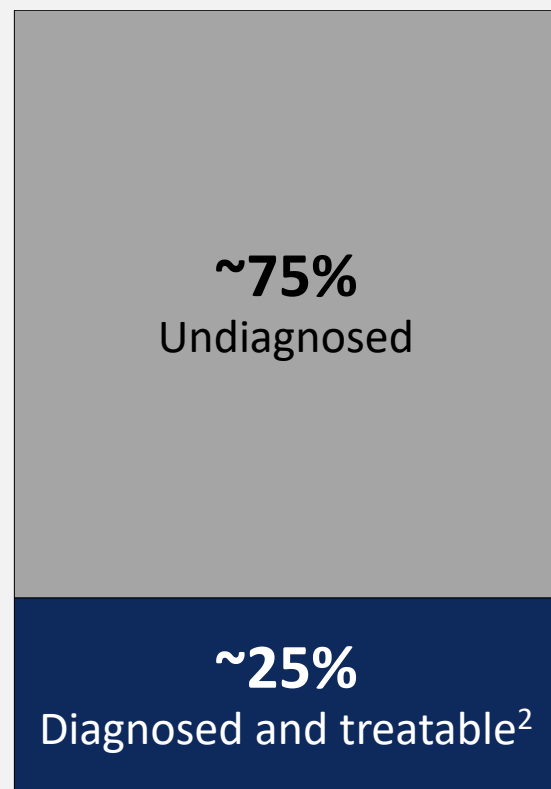
Meta-analysis of published ADH1 case reports



ADH1 market opportunity is growing as genetic diagnoses increase

Estimated US ADH1 population

12,000-13,000 variant carriers¹



2019



\$750M+ worldwide revenue opportunity

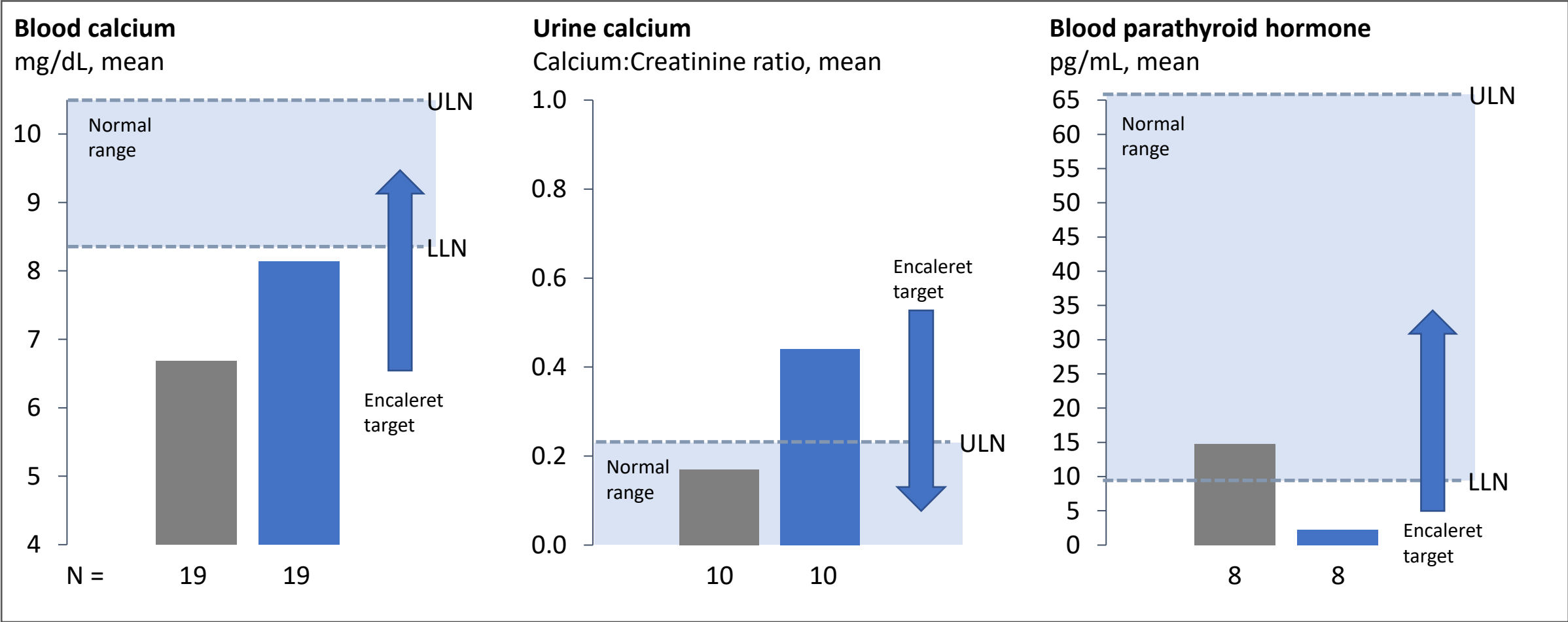
Potential upside if diagnosis rates improve due to:

- Increasing genetic testing, including BridgeBio-sponsored program
- Increasing disease awareness and available targeted therapy

Calcium and active Vitamin D supplementation is ineffective at normalizing blood calcium, urine calcium, and blood PTH in ADH1 patients

Summary of key disease measures in ADH1 patients with and without supplementation

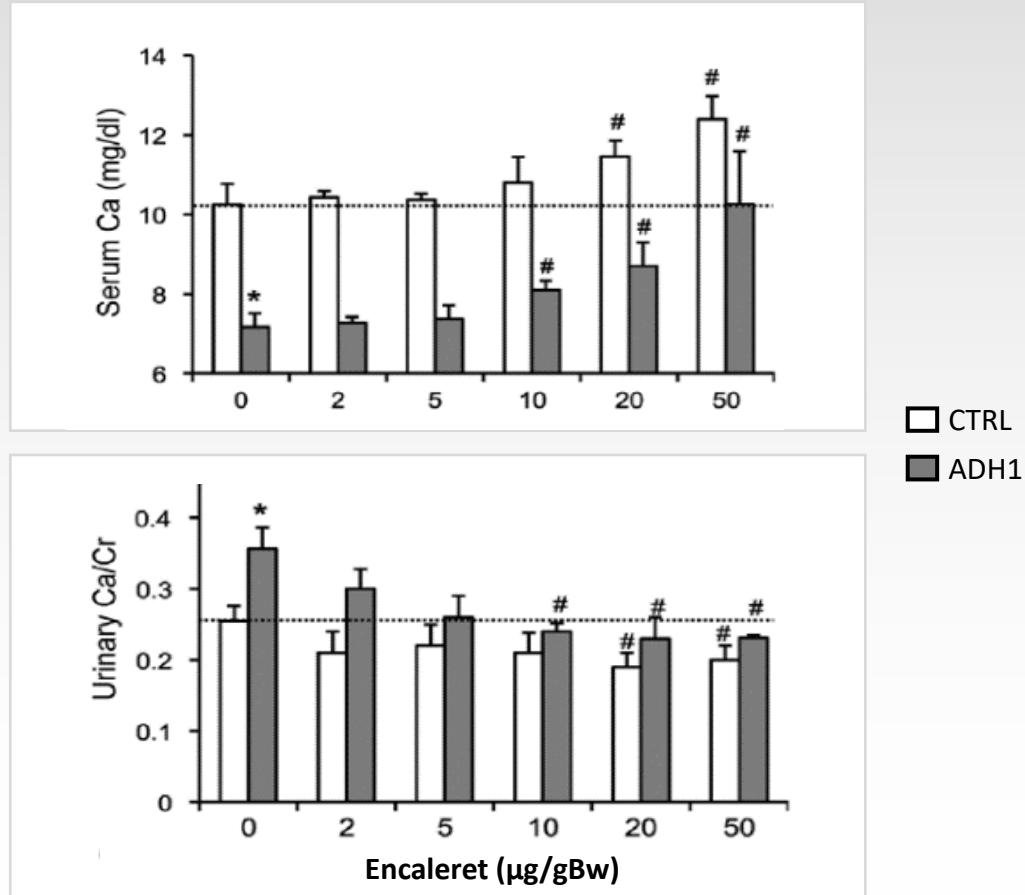
Without supplementation
With supplementation



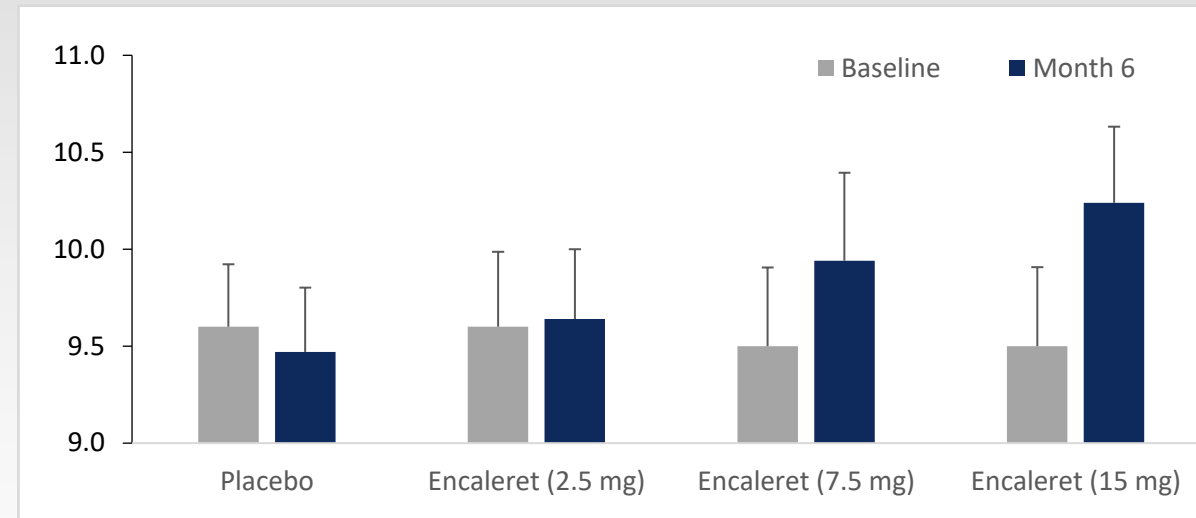
ULN = upper limit of normal, LLN = lower limit of normal
Source: Pearce et al. Clin Endocrinol (Oxf).1996. PTH values reported as below detection limit or undetectable were recorded as "0"

Encaleret has demonstrated proof of mechanism in mouse model of ADH1 and in patients with osteoporosis

Encaleret normalized serum and urine calcium in a mouse model of ADH1¹

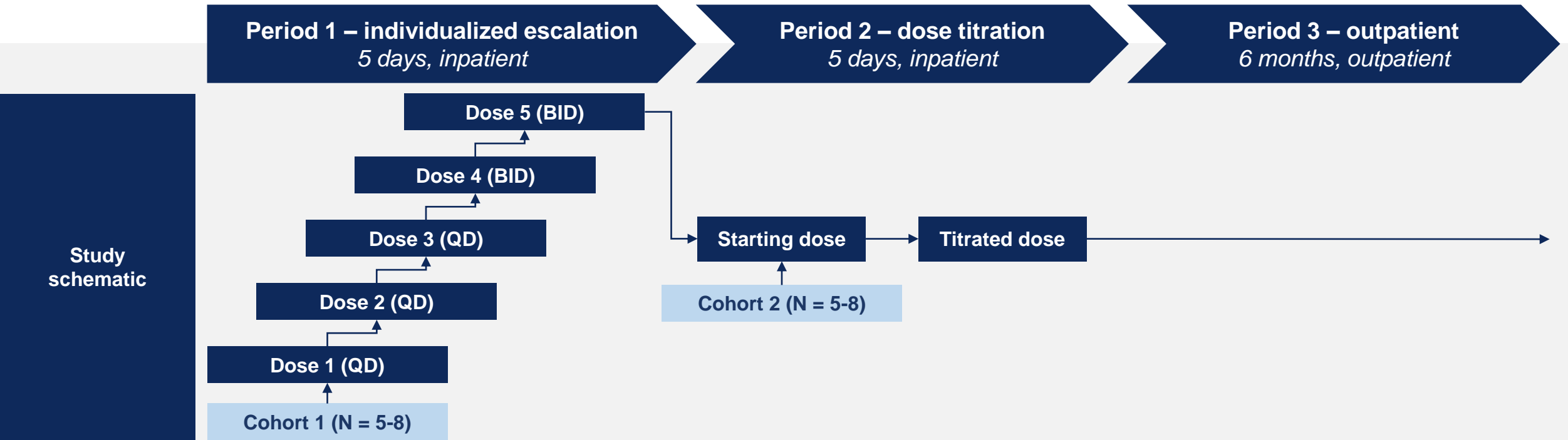


Encaleret was well-tolerated and increased serum calcium in clinical trials in patients with osteoporosis²



Hypercalcemia was dose-limiting safety concern in osteoporosis program (>1,200 participants); increasing serum calcium levels is target effect in ADH1

Phase 2, open-label dose-ranging study is evaluating safety, tolerability, and efficacy of encaleret in ADH1 patients



Potential proof-of-concept in Period 1 based on key objectives:

- Safety and tolerability
- Blood calcium concentration
- Urine calcium concentration
- Intact parathyroid hormone concentration

Additional measures

- Serum 1,25(OH)₂-vitamin D
- Serum Mg, Pi, Na, K, Cr, cAMP, citrate
- Bone turnover markers (serum collagen C-telopeptide, serum procollagen Type 1 N-propeptide)



Acoramidis (AG10) for transthyretin (TTR) amyloidosis (ATTR)

ATTR overview



Prevalence

400,000+ worldwide,
largely undiagnosed today



Genetic driver

Destabilizing TTR
variants or factors of
aging, leading to
amyloid accumulation



Pathophysiology

Systemic disease most
commonly presenting as
cardiomyopathy or peripheral
neuropathy

Features of a potential best-in-class medicine for ATTR



**Near-complete
stabilization of TTR,**
preventing the formation
of amyloid deposits

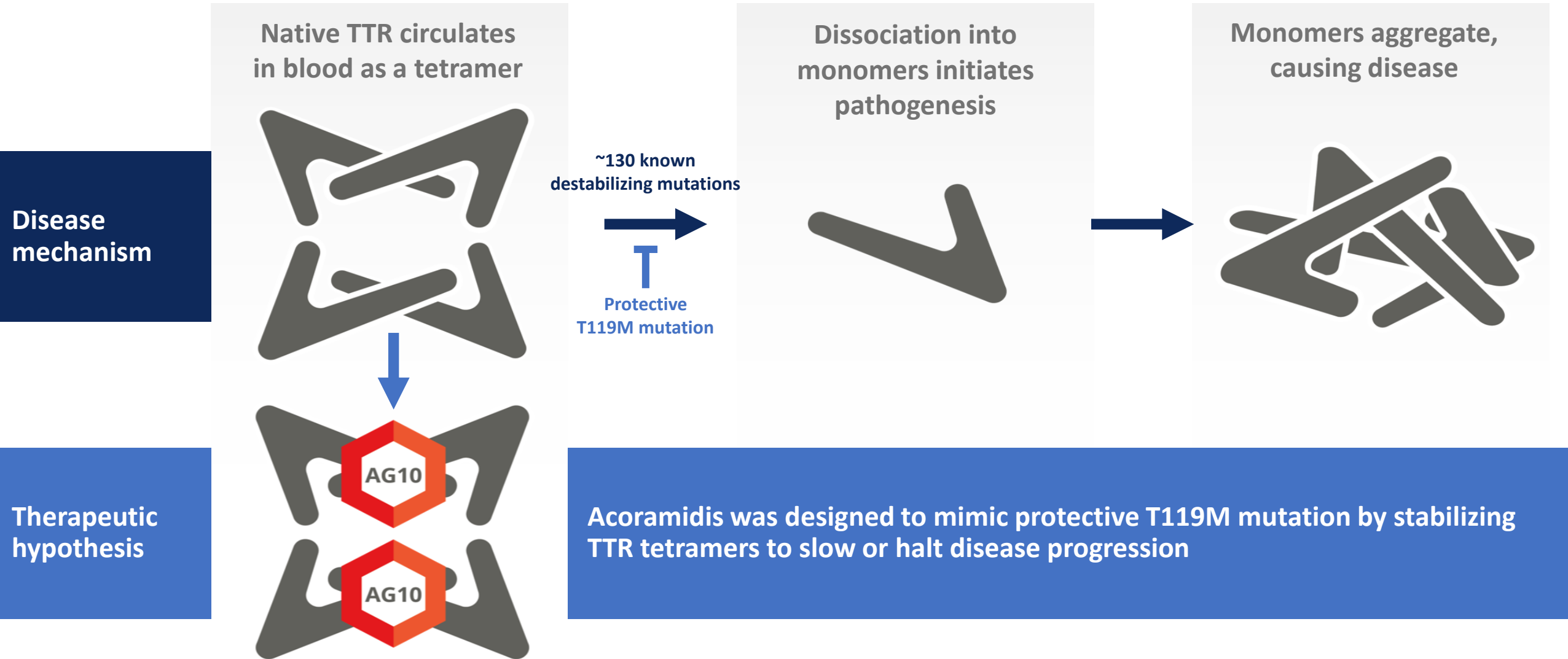


Preserve TTR tetramer,
which has known beneficial
roles and is highly
evolutionarily conserved



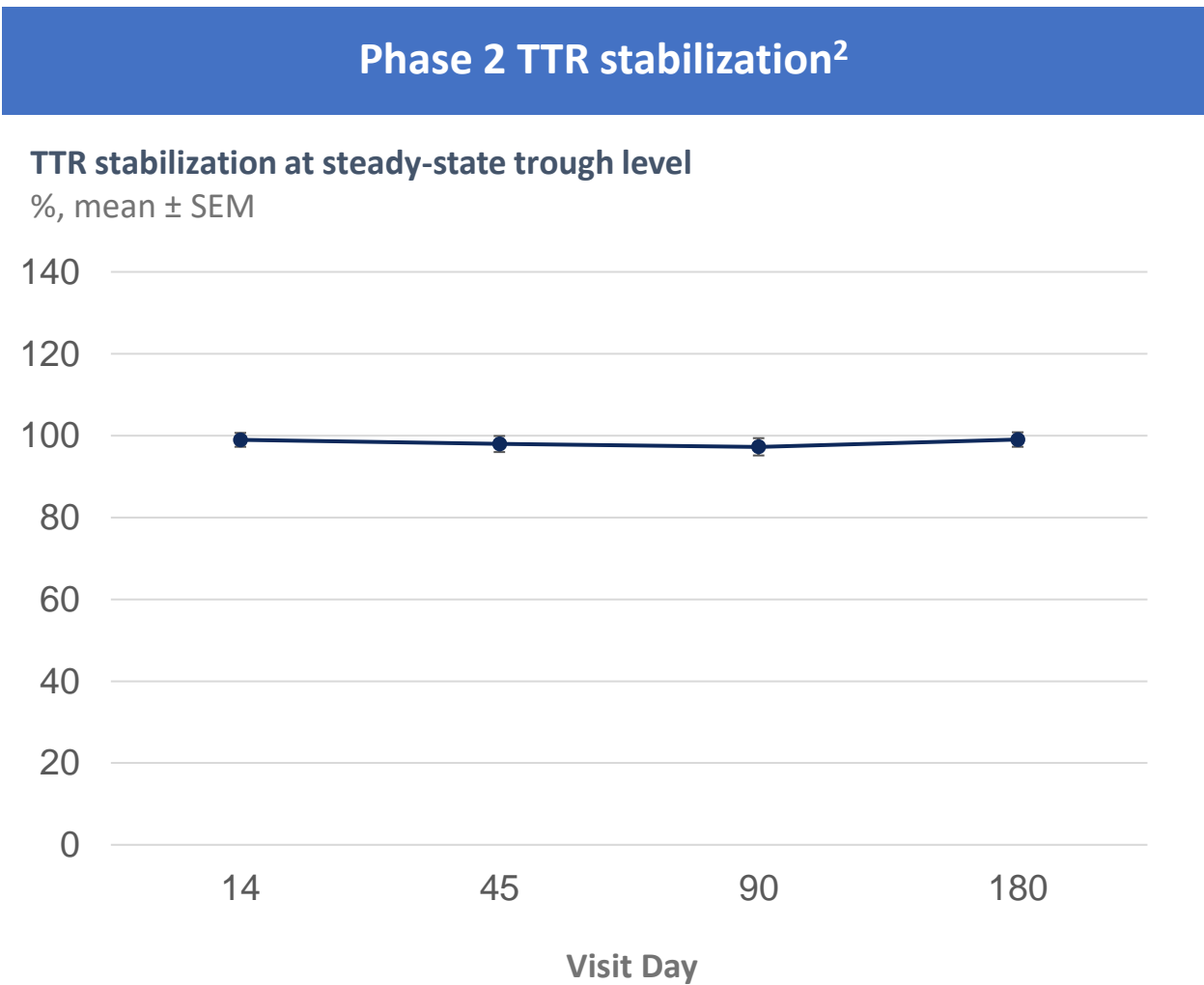
Oral dosing, a convenient and
flexible solution for ATTR
patients and their families

Acoramidis was designed to treat ATTR at its source



Acoramidis has been well-tolerated and demonstrated near-complete TTR stabilization in pre-clinical, Ph1, and Ph2 studies

Phase 2 safety summary ¹		
	Placebo N = 17	Acoramidis (pooled doses) N = 32
Any Adverse Event	15 (88%)	21 (66%)
Mild	6 (35%)	11 (34%)
Moderate	8 (47%)	9 (28%)
Severe	1 (6%)	1 (3%)
Any Serious Adverse Event	2 (12%)	1 (3%)
AF and CHF	1 (6%) ¹	0
Leg cellulitis	1 (6%)	0
Dyspnea	0	1 (3%)

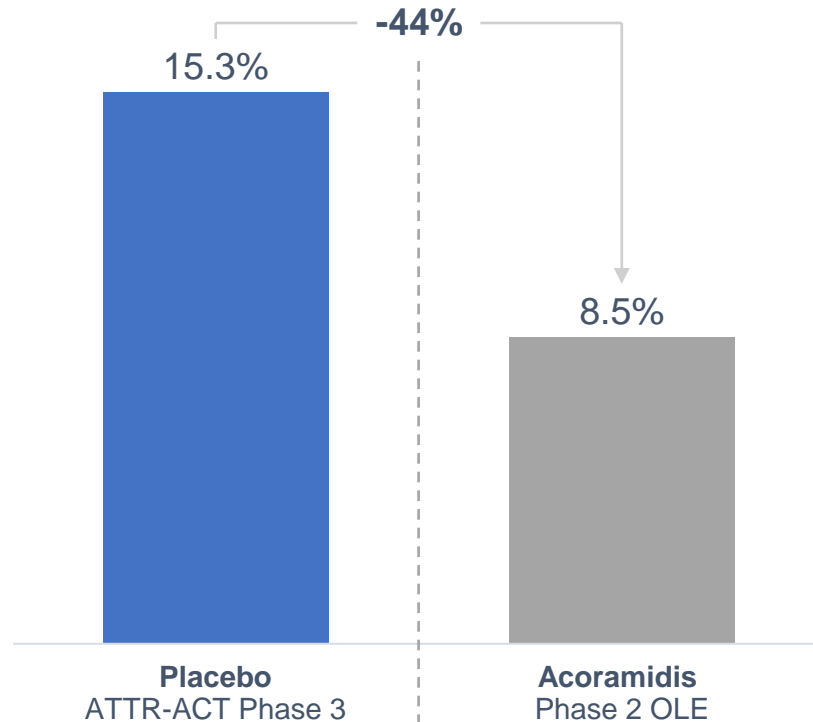


1 Judge, D.P. et al. JACC Vol. 74, No. 3, 2019:285 – 95

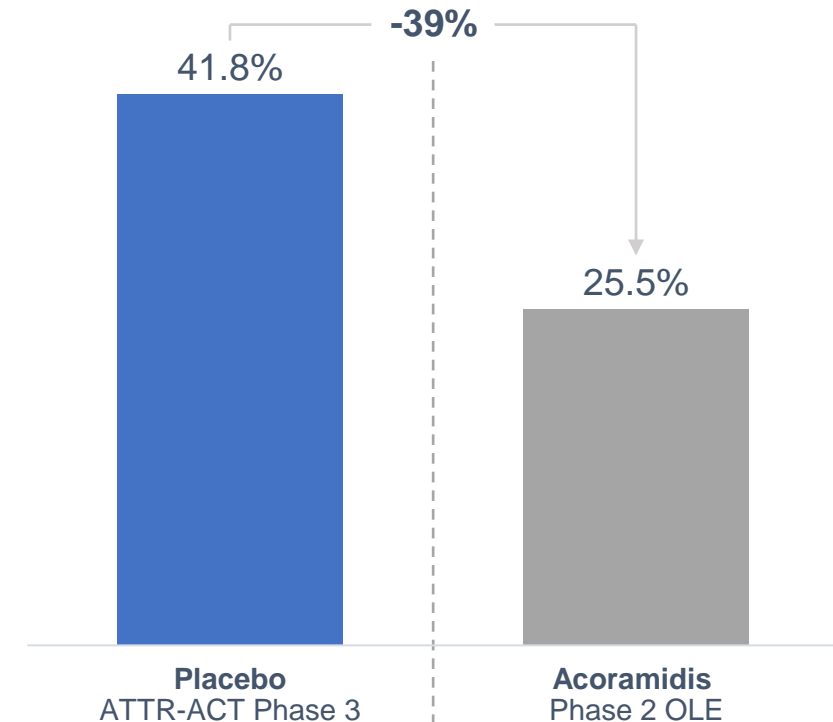
2 Judge, D.P. et al. American Heart Association 2019

Deaths and CV hospitalizations reported in acoramidis Phase 2 OLE were lower than in placebo-treated ATTR-ACT participants

All-cause mortality at 15 months Participants died or receiving transplant (%)



Cardiovascular hospitalizations at 15 months Participants with ≥1 CV hospitalization (%)



¹ Based on routine adverse event reporting

Note: These data are based on a cross-trial comparison and not a randomized clinical trial. As a result, the values shown may not be directly comparable

Source: Judge, DP et al. American Heart Association Scientific Sessions 2019

ATTRibute-CM will provide 12-month functional outcome data and 30-month mortality and CV hospitalization data

Key inclusion criteria

- Subjects with diagnosed ATTR-CM (WT or mutant)
- NYHA Class I-III
- ATTR-positive biopsy or ^{99m}Tc scan
- Light chain amyloidosis excluded if diagnosis by ^{99m}Tc

Screening and randomization

12-month primary endpoint:
Change in 6MWD

30-month primary endpoint:
Mortality and CV hospitalizations

800 mg acoramidis twice daily

Target N ~ 340

Placebo twice daily

Target N ~170

800 mg
acoramidis
twice daily

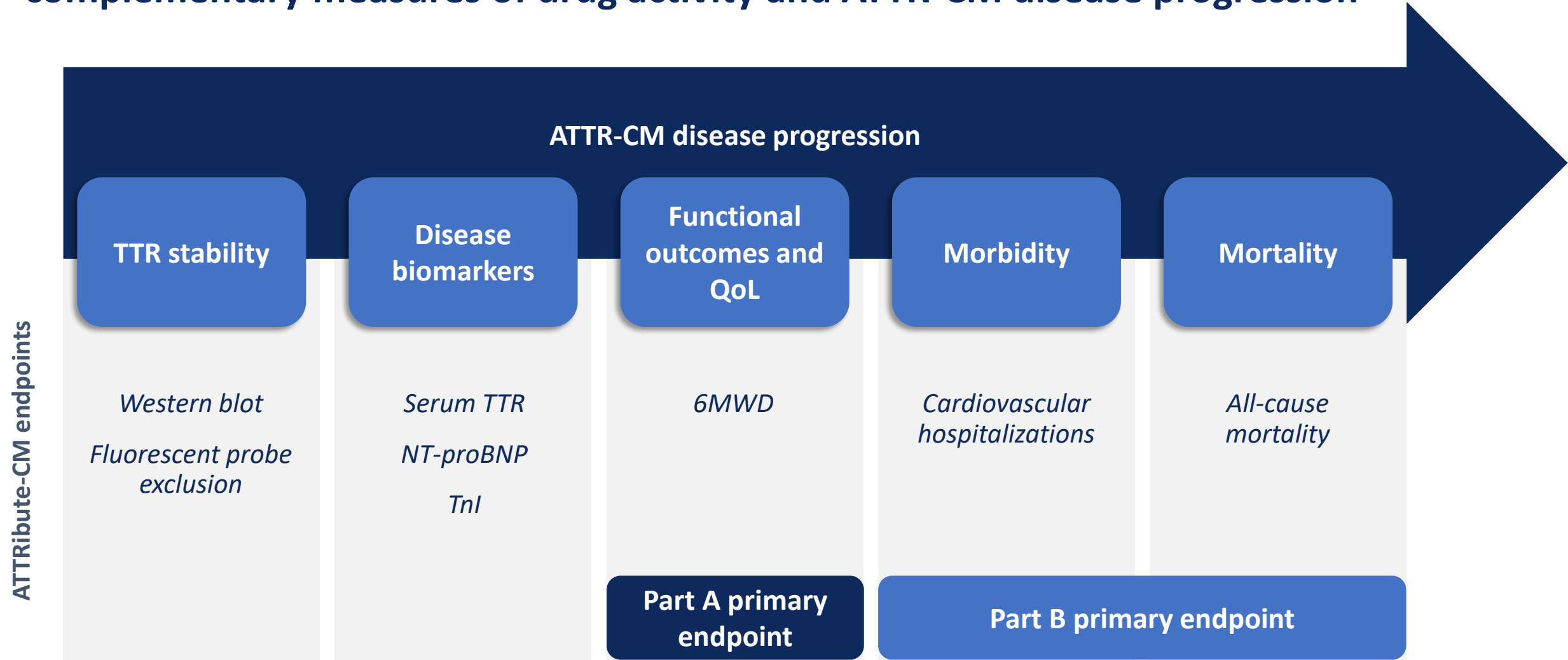
Part A

Part B
Tafamidis usage allowed

Open label extension

6MWD = Six-minute walk distance; NYHA = New York Heart Association;
^{99m}Tc = Technetium labeled pyrophosphate (PYP) or bisphosphonate (e.g., DPD);
CV = cardiovascular-related

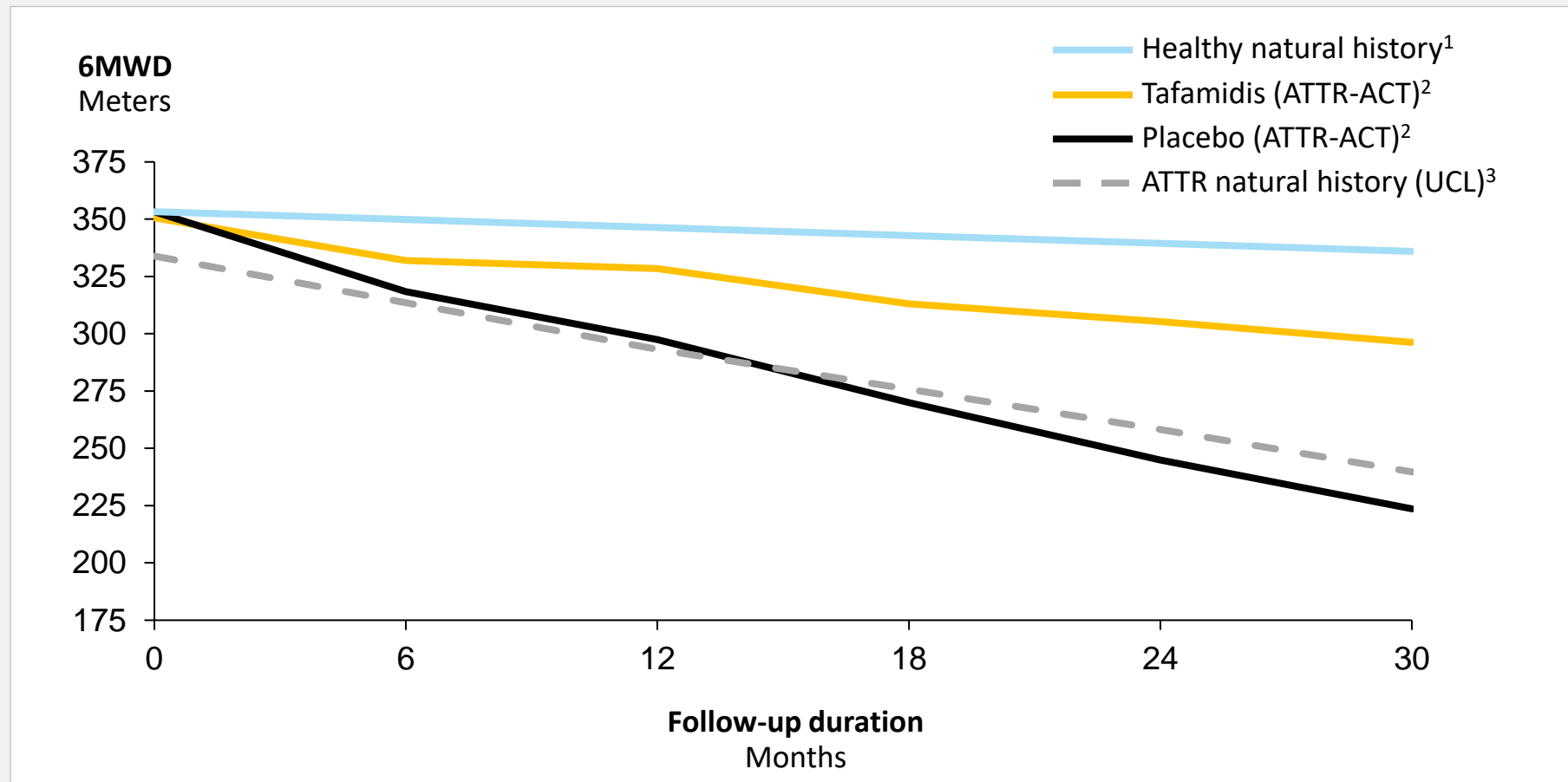
ATTRibute-CM is designed to evaluate safety and efficacy of acoramidis across complementary measures of drug activity and ATTR-CM disease progression



NT-proBNP = N-terminal pro b-type natriuretic peptide; TnI = Troponin I; 6MWD = Six-minute walk distance
QoL = Quality of life

Rapid functional decline in untreated ATTR-CM patients provides opportunity to demonstrate robust clinical benefit

Summary of 6MWD data in ATTR-CM and healthy cohorts



Optimal profile for tafamidis would markedly slow or halt decline in 6MWD in trial participants

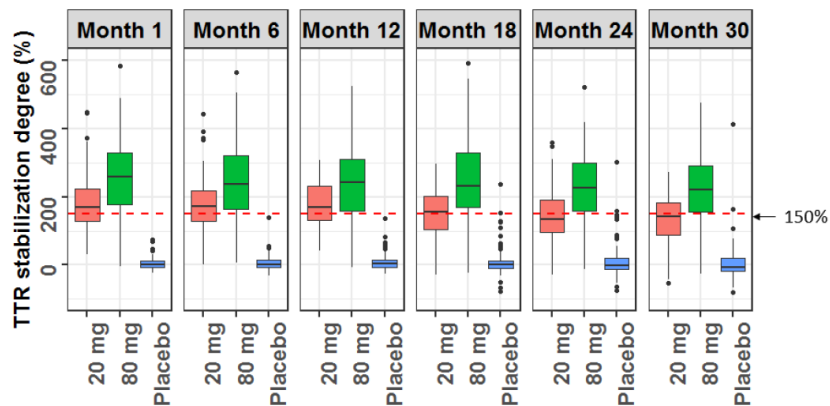
1. Enright, P.L. et al. Chest 2003. N = 3333 healthy elderly adults, baseline set to match ATTR-ACT placebo group
2. Maurer, M.S. et al. NEJM 2018. N = 264 (tafamidis), N = 177 (placebo) ATTR-CM trial participants
3. Lane, T.L. et al. Circulation 2019. N = 1034 ATTR-CM patients

Higher dose of tafamidis demonstrated increased TTR stabilization and greater clinical benefit in ATTR-ACT + LTE

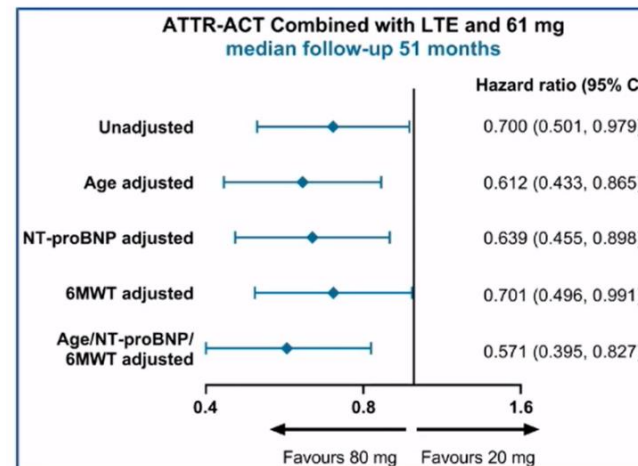
Phase 3 ATTR-ACT study tested two doses of tafamidis (20 mg & 80 mg) vs. placebo

- In an analysis of ATTR-ACT combined with long-term extension (LTE), benefit of tafamidis 80 mg vs. 20 mg was evident on all-cause mortality¹
- At baseline, ATTR-ACT participants treated with 80 mg of tafamidis were older and had more severe evidence of disease than those treated with 20 mg of tafamidis¹
- Participants receiving 80 mg of tafamidis (vs. 20 mg) exhibited greater TTR stabilization²

TTR stabilization²



All-cause mortality¹



**Increased levels
of TTR
stabilization may
translate to
improved clinical
outcomes in
ATTR-CM**

1. Damy, T., ESC Heart Failure Association Discoveries 2020. "The Tafamidis in Transthyretin Cardiomyopathy Clinical Trial."

2. FDA CDER Clinical Pharmacology and Biopharmaceutics, Clinical Review (Vyndaqel/Vyndamax), 2019; Fourfold increase in tafamidis dose did not lead to a fourfold increase in TTR stabilization due to non-linear pharmacokinetics

Ongoing and planned studies of acoramidis aim to continually expand clinical evidence and addressable patient population

ATTRibute-CM Phase 3 study enrolled 632 participants and is on track for topline data in late 2021 or early 2022

2021

ATTRibute^{CM}

ATTR-CM
WT and hereditary
Functional outcomes

2023

ATTRibute^{CM}

ATTR-CM
WT and hereditary
Functional outcomes
+
Composite mortality and morbidity

2024

ATTRibute^{PN}

ATTR-PN
Hereditary
Functional outcomes

ATTRibute^{PN}

ATTR-PN
Hereditary
Functional outcomes

2025+

Prevention in high risk populations
Head-to-head comparisons



Claudia,
child with achondroplasia

Low-dose FGFR inhibitor (infigratinib) for achondroplasia

Achondroplasia overview



Prevalence

55,000 (US+EU) –
one of the most common
genetic conditions



Genetic driver

FGFR3 activation



Pathophysiology

Up-regulation of STAT1 and MAPK
in the growth plate cause cranial,
spinal, and stature symptoms

Features of a potential best-in-class medicine for achondroplasia



Direct targeting of FGFR3

and normalization of both
STAT1 and MAPK
signaling pathways



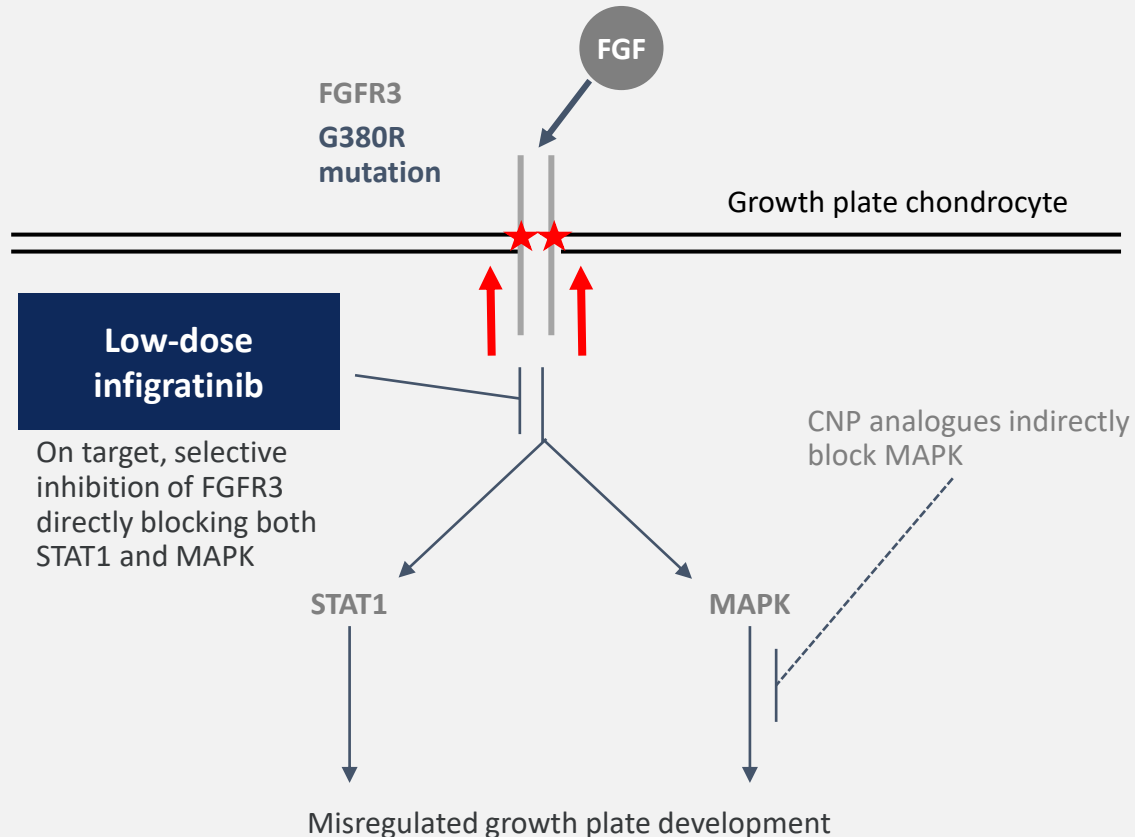
Potential to address all

drivers of symptoms,
including cranial, spinal
and stature issues



Oral dosing, the most
convenient solution for
children with achondroplasia
and their families

Potential best-in-class approach targeting achondroplasia directly at its genetic source



ACH FGFR3 gain-of-function mutation causes:

- 2-3x over-activation of the receptor
- Up-regulation of downstream pathways STAT1 and MAPK
- Aberrant growth plate development, which causes cranial, spinal, and stature symptoms

Low-dose infgratinib has the potential to:

- Directly inhibit the causal gain-of-function mutation in FGFR3
- Normalize both the STAT1 and MAPK signaling pathways
- Reverse all key drivers of symptoms

Low-dose infigratinib improves all the key drivers of clinical symptomology in validated ACH mouse model

1 Cranial bone issues

17%

increase in
FM area

6%

increase in AP
skull length

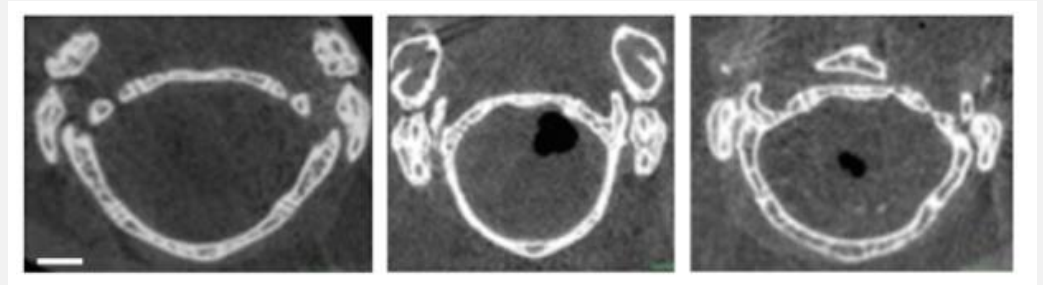


May lead to **decrease in foramen magnum stenosis** and fewer surgeries

FGFR3 WT
No treatment

FGFR3^{Y367C/+}
No treatment

FGFR3^{Y367C/+}
Infigratinib tx



2 Disorders of the spine

12%

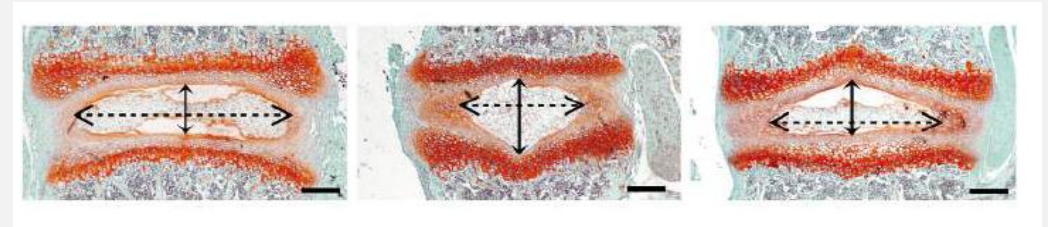
increase in
L4-L6 length

73%

increase in
disc width



May lead to **decrease in spinal stenosis**, possibly **reducing need for surgery**



3 Disproportionate short stature

21%

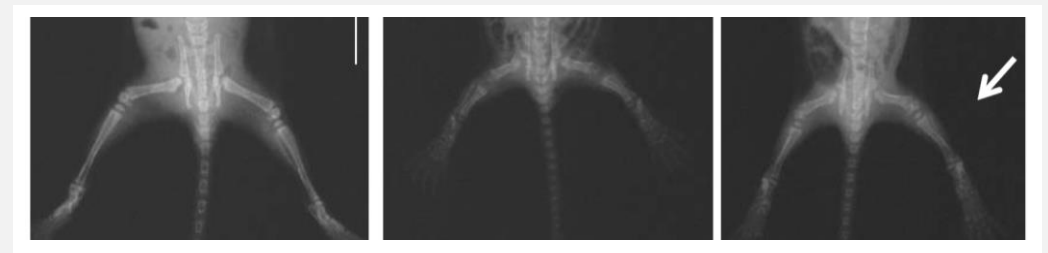
increase in
femur length

33%

increase in
tibia length





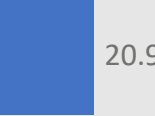
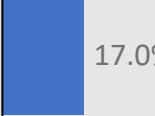
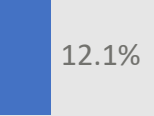
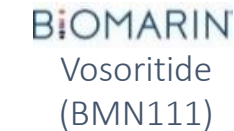








May lead to **increased stature and proportionality**



Source: Komla-Ebri et al. J Clin Inv 2016

Note: percent increase compared to vehicle treated FGFR3^{Y367C/+} mouse, infigratinib treatment with 2mg/kg subcutaneous dose

Low-dose infigratinib showed potential best in-class preclinical profile in validated achondroplasia mouse model

Company/ Asset	MOA	Route	Status	Mouse model	Tibia length	Femur	Foramen magnum area	L4-L6 height
 Infigratinib	Selective FGFR1-3i	Oral	Ph2	FGFR3 ^{Y367C/+}	 32.6%	 20.9%	 17.0%	 12.1%
 Vosoritide (BMN111)	CNP analogue	Daily SQ	Pivotal (NDA filed)	FGFR3 ^{Y367C/+}	 6.6%	 5.2%	<div>No known publicly available data</div>	 3.3%
 TransCon CNP ¹	CNP analogue	Weekly SQ	Ph2	FGFR3 ^{Y367C/+}	 12.3%			
 Reifercept (TA-46)	FGFR3 decoy	Weekly SQ	Ph2	FGFR3 ^{ACH}	 8.6%	 6.2%		

Preclinical data from infigratinib and other investigational achondroplasia therapies

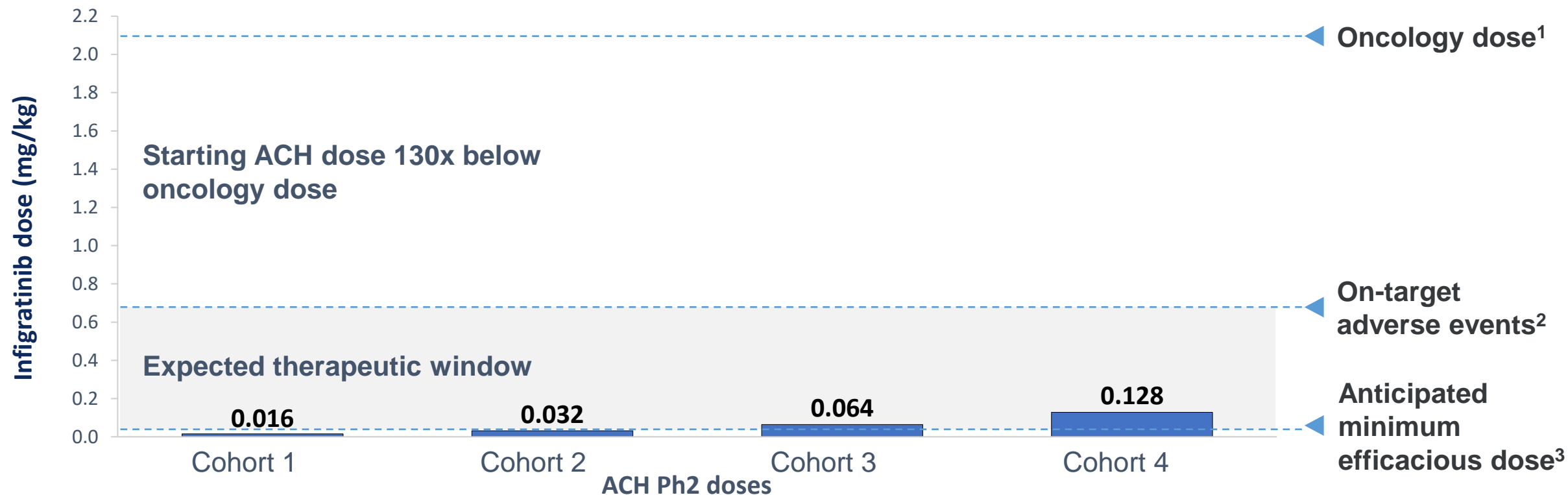
Percent increase compared to non-treated mouse

Source: Komla-Ebri et al. J Clin Inv 2016, Lorget et al. Am J Hum Genet 2012, Garcia et al. Science Trans Med 2013, Breinholt ENDO 2017
Note: subcutaneous doses, percent increase compared to vehicle treated FGFR3^{Y367C/+}, FGFR3^{ACH/+} mouse as noted in “Mouse model” columns
Infigratinib treatment with 2mg/kg subcutaneous dose ¹Based on vosoritide continuous infusion; *Value estimated using Digitizelt.

We have a wide anticipated therapeutic index in achondroplasia

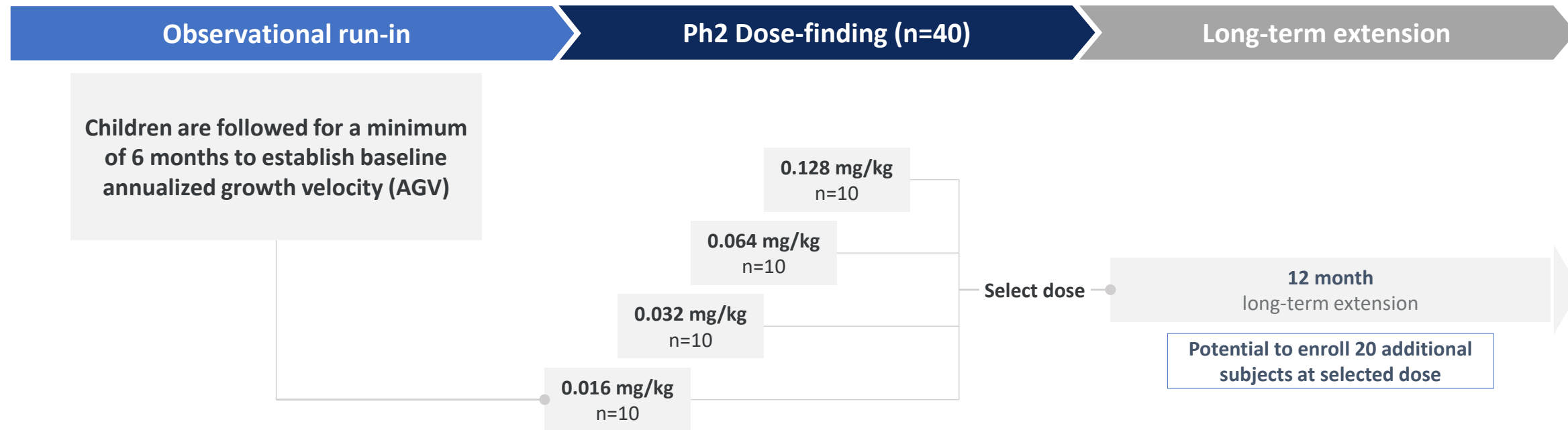
Infigratinib has been tested in >700 humans in our oncology program, providing significant data on PK, tolerability and safety

Most common and dose-limiting side effect is phosphorus elevation (on-target through FGFR1 inhibition), which occurs significantly above our planned achondroplasia doses



¹Based on 125mg dose and 60kg adult; ²Based on estimated TD₅₀ at 40mg and 60kg adult; ³Based on PK modeling and allometric scaling from animal models

The PROPEL clinical program is enrolling with data expected in 2H 2021



Key inclusion criteria

- Children 2.5 – 10 years old
- Clinical and molecular ACH diagnosis

Primary objectives

- Baseline annualized growth velocity (AGV)

Primary objectives

- Identify safe therapeutic dose for expansion / pivotal study
- Safety and tolerability
- Change from baseline in AGV

Primary objectives

- Long-term safety and efficacy

BBP-631: AAV5 gene therapy for congenital adrenal hyperplasia (CAH)

Program overview



Prevalence

75,000 (US+EU) – One of the largest known AAV gene therapy markets



Genetic driver

21-hydroxylase inactivation



Pathophysiology

Inability to produce cortisol causes need for supraphysiologic doses of synthetic steroids, 3x increase in mortality risk, hirsutism, Cushingoid symptoms

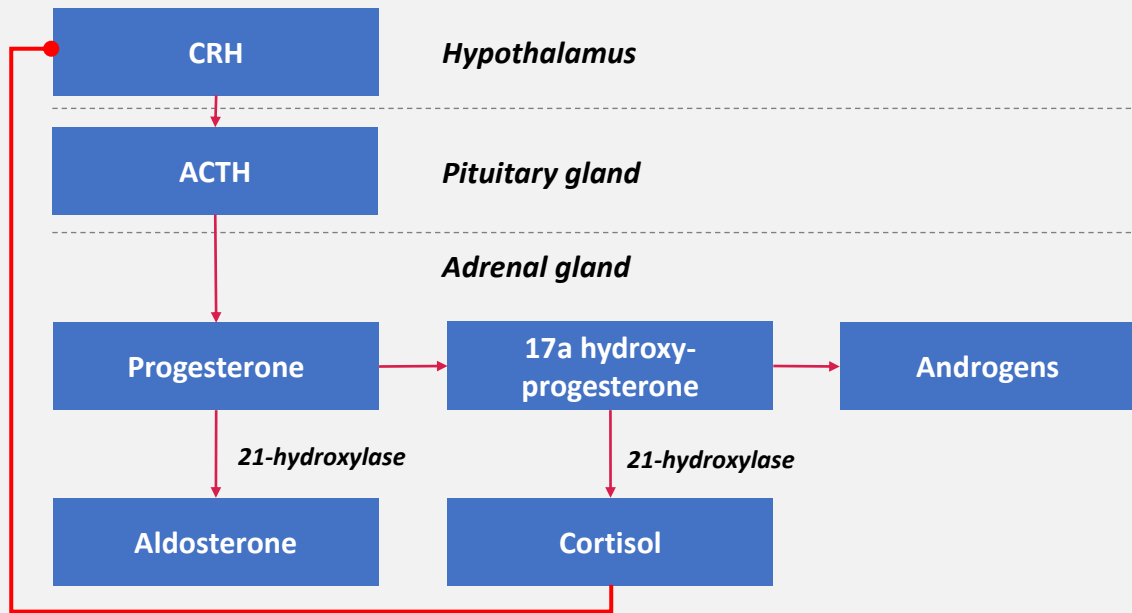
We believe CAH is an ideal indication for AAV gene therapy:

- **Low threshold to correct phenotype**, validated by human clinical genetics (~5-10% of WT enzyme activity)
- **Only approach designed to induce endogenous cortisol and mineralocorticoid production**, potentially allowing steroid withdrawal
- **Durable transgene delivery to the adrenal gland of NHPs** with IV dosing of our construct
- **Next catalyst:** initial data from first-in-human study

Maris,
child with CAH

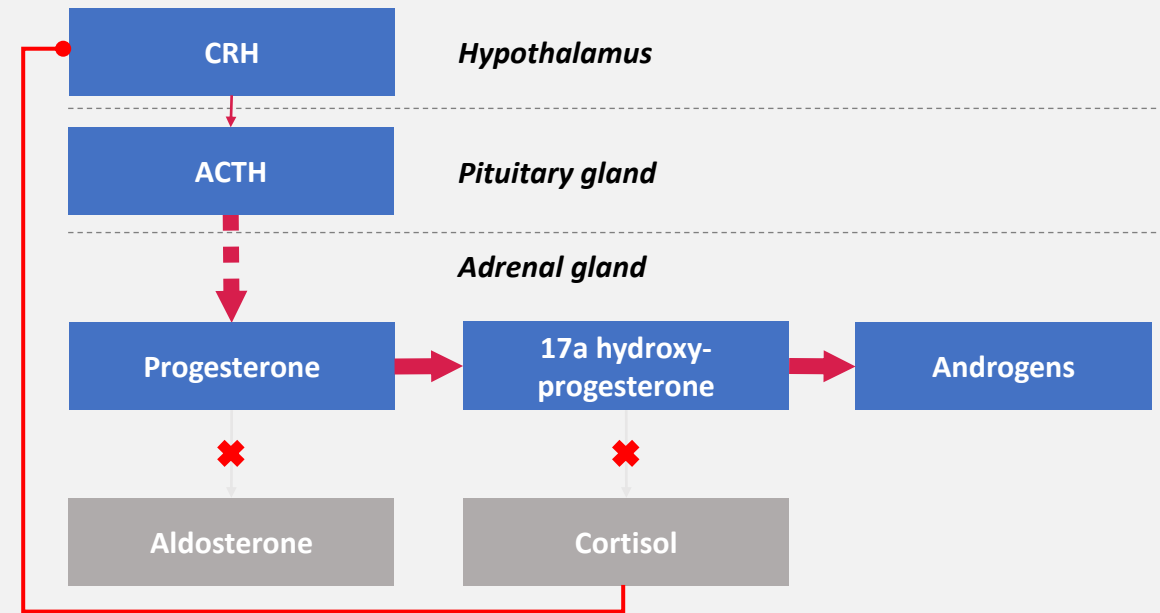
Gene therapy is the only modality designed to treat CAH at its source and allow for production of endogenous cortisol

Healthy Hypothalamic-Pituitary-Adrenal Axis



In a functional HPA system, cortisol and aldosterone are produced as needed by the body. Cortisol serves as a “brake” on the CRF/ACTH system

Hormonal dysregulation with 21OHD; no cortisol “brake” on ACTH, shunting of 17OHP to androgens

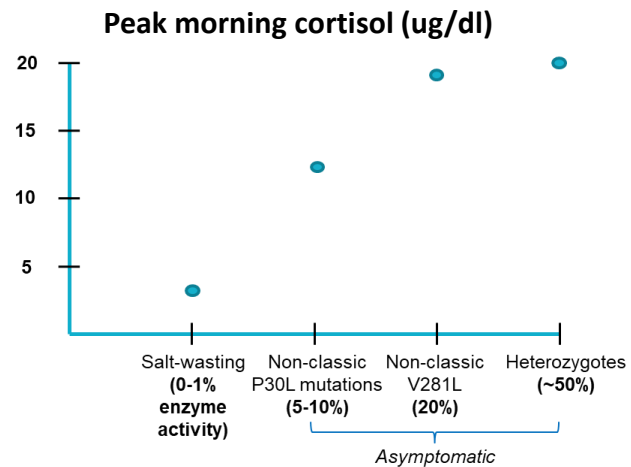


In CAH, cortisol and aldosterone are not able to be produced. The lack of a “cortisol brake” results in buildup of progesterone and 17OHP, leading to an excess of androgen production

CAH patients have 3-4X higher mortality than the general population, and suffer significant morbidity ranging across cardiovascular and metabolic disease, bone disease, infertility, chronic fatigue, and other disorders.

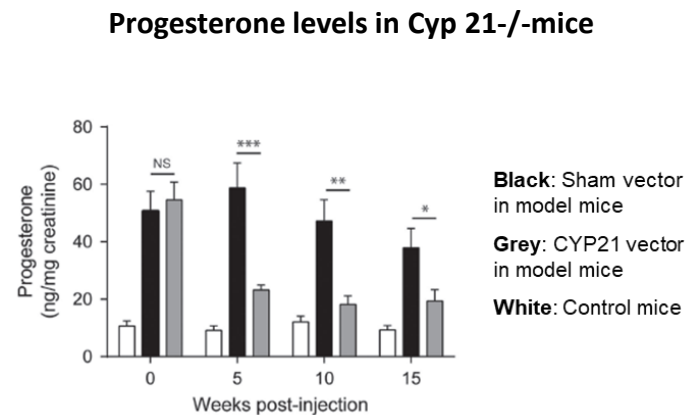
CAH: NHP study showed durable transgene expression; 5-10% of WT enzyme may be sufficient for clinical impact

Genotype-phenotype studies show that >5-10% of enzyme activity results in nonclassical CAH



- Due to the high enzymatic efficiency/selectivity of 21-OHase, **only a small amount of enzyme is required to rescue the phenotype**

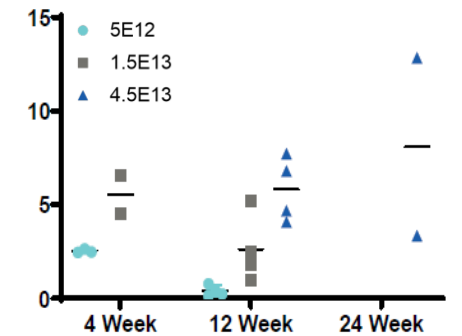
Mouse studies show a VGC of only 0.13 at 18 wks was sufficient for phenotypic correction



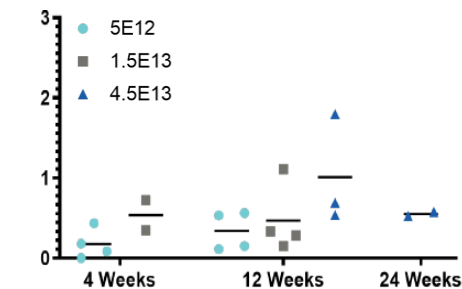
- At 15 weeks in treated mice, **progesterone** (the key substrate of 21OHase in mice) was **significantly reduced vs untreated mice**

NHP studies show sustained VGC and RNA out to 6 months

VG Copies per Cell (DNA)



hCYP21A2/Ywhaz (RNA)

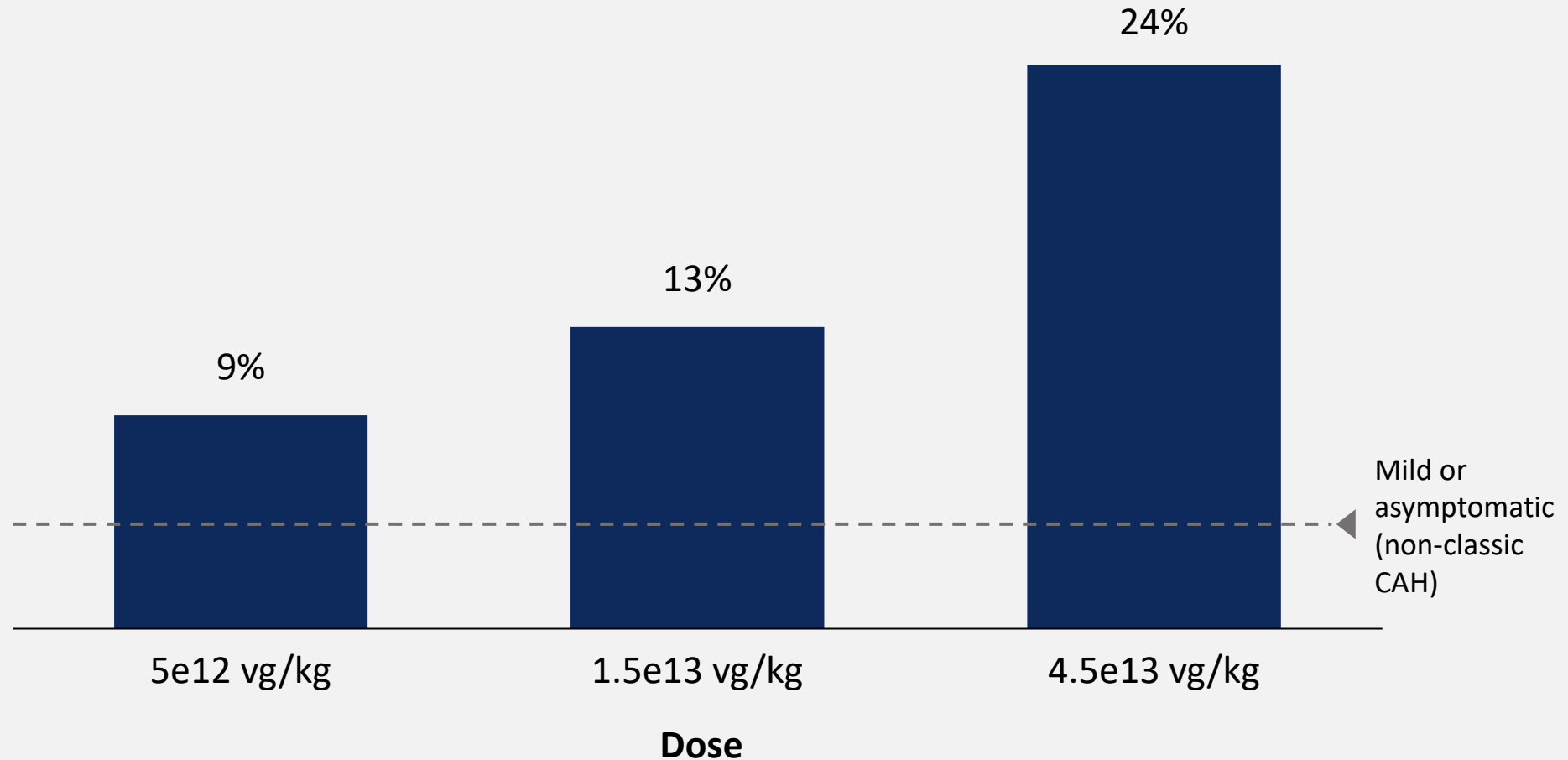


- Transgene expression is dose-dependent and stable out at 24 wks
- We can durably transduce the NHP adrenal gland with our construct at >20x the vector required to correct the CAH phenotype in mice**

NHP protein data using mass spec methods suggests potentially therapeutic levels of 21-hydroxylase enzyme

Human 21-hydroxylase protein as a % of NHP 21-hydroxylase protein (Mass Spec quantification)

- We have developed mass-spec methods to quantify protein expression by identifying differential peptides between human and NHP 21-OH
- These data suggest dose-dependent enzyme expression in the adrenal cortex from 9%-24% of WT levels
- Genotype-phenotype relationship suggests as little as 5% of WT enzyme activity is associated with the mild/asymptomatic non-classic form of CAH





BridgeBio oncology research

World-class oncology team drives our discovery and development

Eli Wallace

CSO Oncology Research



Pedro Beltran

SVP Oncology



Frank McCormick

Chairman of Oncology



Richard Scheller

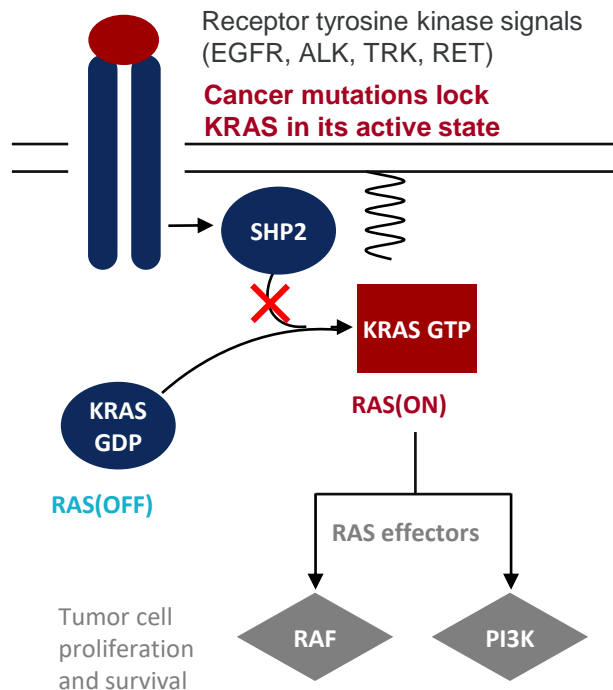
Chairman of R&D



Basia, pancreatic cancer
patient (>90% KRAS-driven)

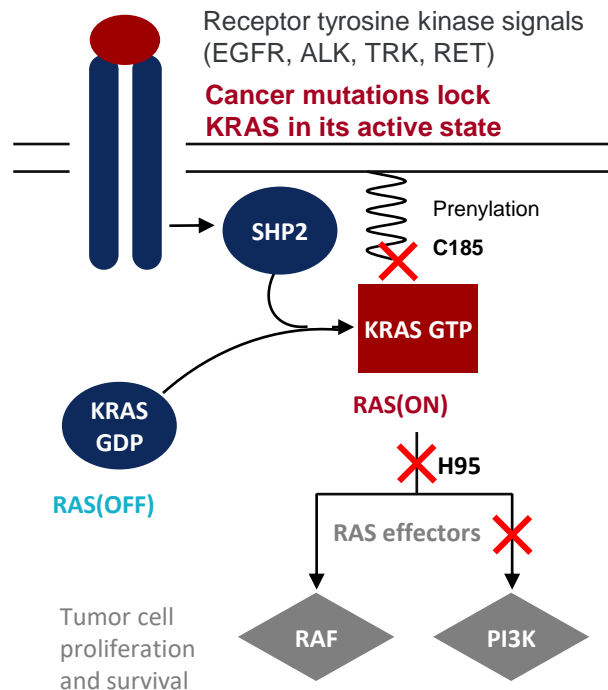
Three disclosed oncology research targets

SHP2 (BBP-398)



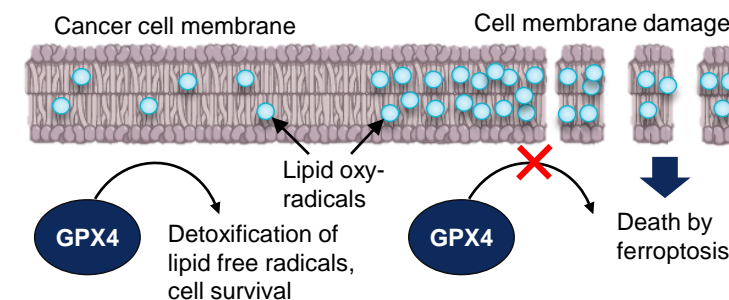
- Potential best in class oral compound
 - Optimized safety, PK and PD profile
 - Maximizes combination therapy potential
- First-in-human study initiated 4Q20

KRAS



- Multiple unexploited sites
- Comprehensive pan-mutant targeting approaches

GPX4



- Potential first in class compound for novel cancer target
- In vivo monotherapy activity and combo potential

Partnerships afford us exceptional collaborators and resources



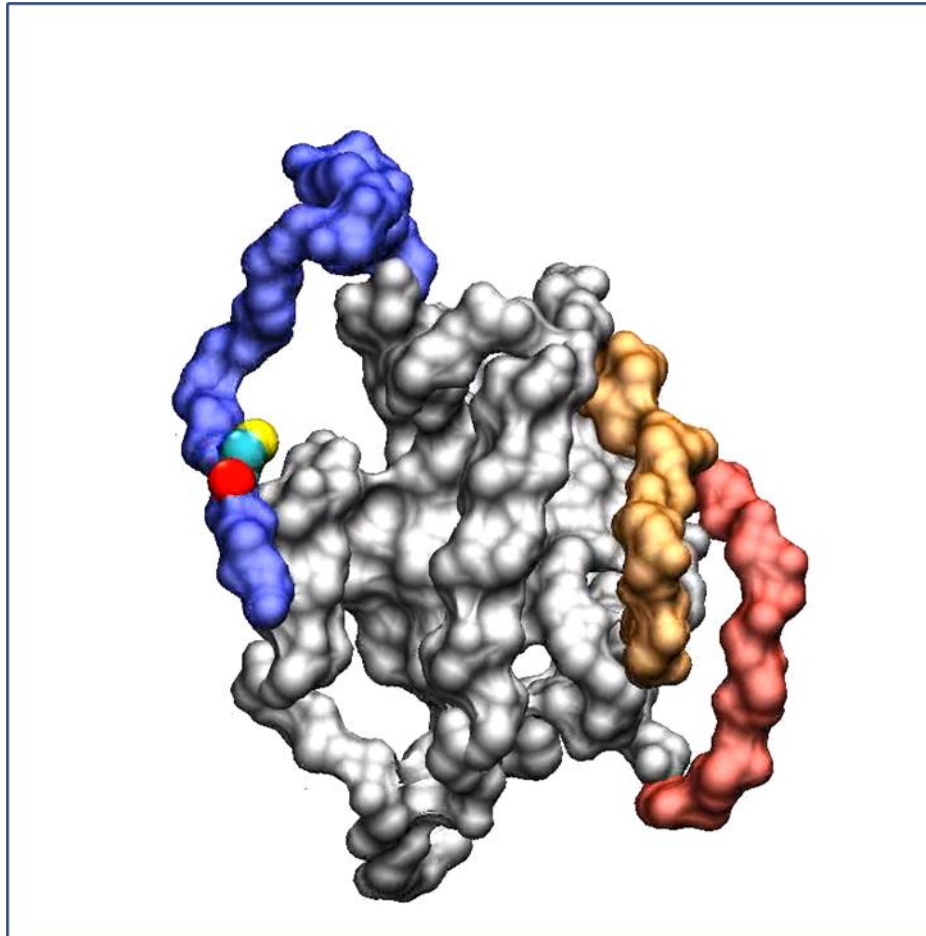
- Partnership with the National RAS Initiative, including 60 of the world's **foremost academic RAS researchers**
- Cutting edge RAS **structural biology expertise**
- Utilization of **cutting-edge instrumentation and techniques**, as well as the **expertise** to lead experiments



- Home to Sierra: the **world's 3rd fastest computing system**
- Enables **multi-microsecond molecular dynamics simulations** of protein complexes, and highly efficient *in silico* **docking simulations**
- This computing power, combined with RAS structural biology expertise at the NCI, delivers **unique insights that fuel our drug design**

Crystal structure enables a static understanding of the target ...

KRAS4b model based on crystal



G-domain
G-domain switch I

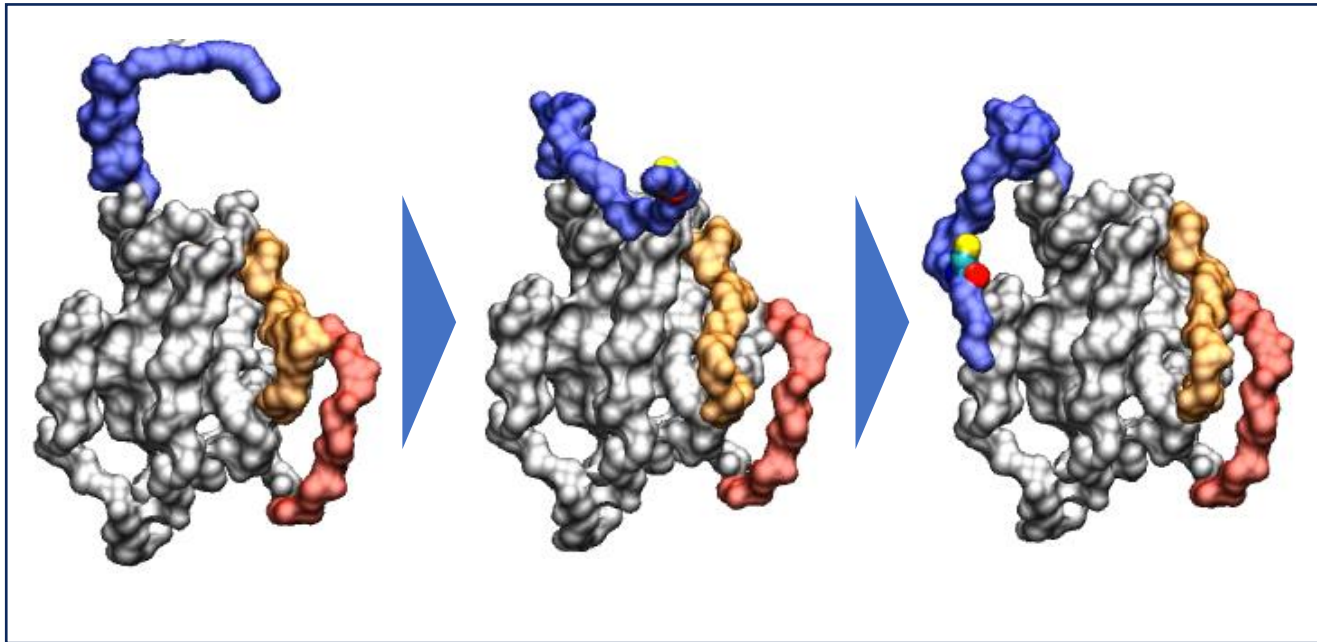
G-domain switch II
Hypervariable region

One therapeutic approach is to inhibit KRAS4b **membrane localization** by targeting **hypervariable region**

Static model reveals only a **subset of potential binding sites** for pharmacological compounds

... whereas molecular dynamics simulation reveals transient conformations and interactions

KRAS4b simulation



G-domain
G-domain switch I

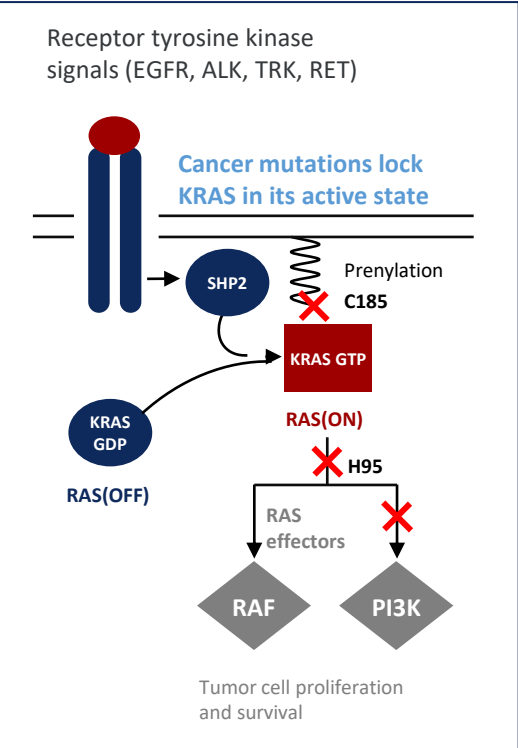
G-domain switch II
Hypervariable region

Reveals possible KRAS4b **HVR transient localization to G-domain**

Elucidates potential transient druggable pocket where **compounds could react covalently with C185**

Enables *in silico* SAR to **inhibit KRAS4b membrane localization**

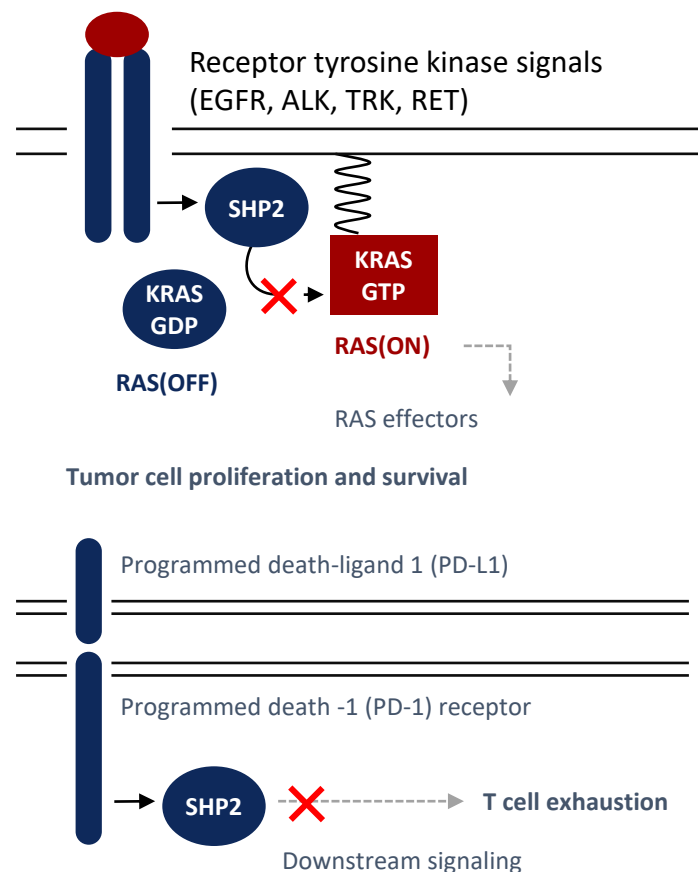
KRAS: multiple shots on goal with our pan-mutant inhibitor programs – each with a unique MOA targeting a novel pocket

KRAS pathway in cancer	Program	MOA	Targets KRAS GTP	Pan-mutant	Crystal structure	Molecular Dynamics
 <p>Receptor tyrosine kinase signals (EGFR, ALK, TRK, RET)</p> <p>Cancer mutations lock KRAS in its active state</p> <p>SHP2</p> <p>Prenylation C185</p> <p>KRAS GTP</p> <p>RAS(ON)</p> <p>H95</p> <p>RAS effectors</p> <p>RAF</p> <p>PI3K</p> <p>Tumor cell proliferation and survival</p>	Program 1: H95 targeting	<ul style="list-style-type: none"> Directly binds activated KRAS through H95 Inhibits KRAS from signaling through effectors 	✓	✓	✓	✓
	Program 2: PI3K effector blocking	<ul style="list-style-type: none"> Blocks specific interaction between KRAS and PI3Ka Blocks PI3K / AKT effector signaling 	✓	✓	✓	✓
	Program 3: C185 targeting	<ul style="list-style-type: none"> Blocks KRAS from tethering Blocks conversion of inactive KRAS GDP to active KRAS GTP 	✓	✓		✓

Our programs are designed to address all KRAS driver mutations, which occur in >30% of all cancers

SHP2: Our compound shows best-in-class potential

Our SHP2i blocks downstream MAPK signaling and abrogates T cell exhaustion



We believe BBP-398 has the ideal properties for combination with a multitude of other therapeutic classes

Human half life: ~15 hours¹

- Allows for recovery above EC50 and reduced MAPK-driven tox

SHP2i combination potential

US + EU incidence, '000s

Potentially differentiated safety profile for combination therapy

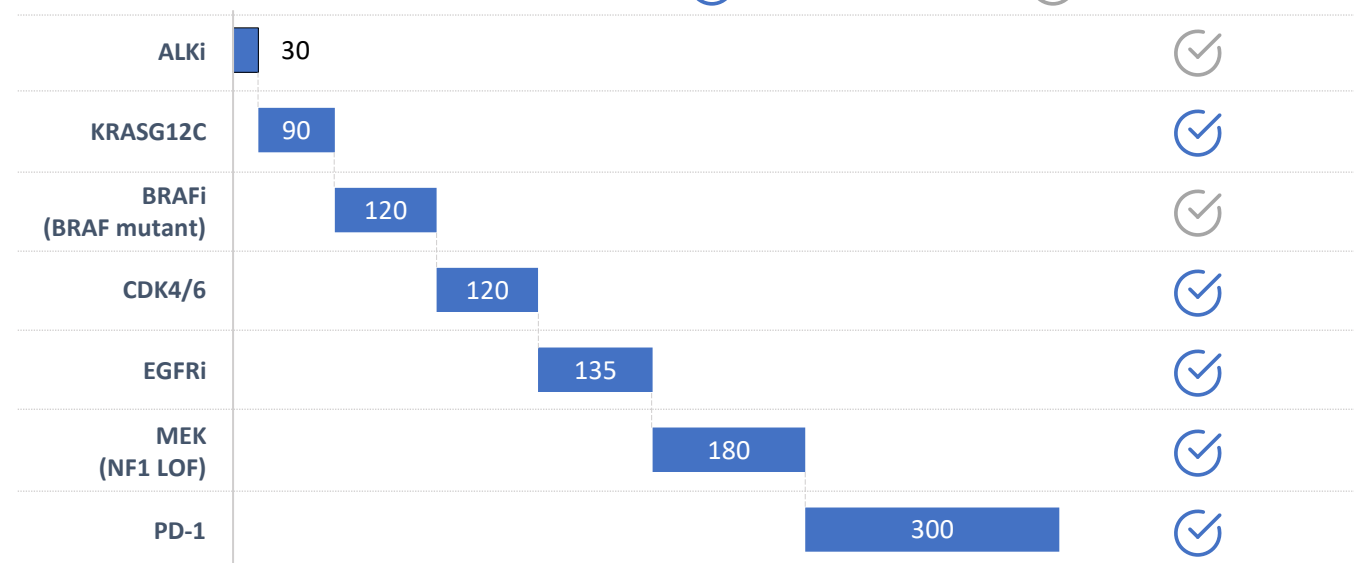
- hERG IC50 (μM)*: >100: No QT prolongation

Supporting evidence²

Peclin data:

✓ BBIO SHP2i

✓ Other SHP2i



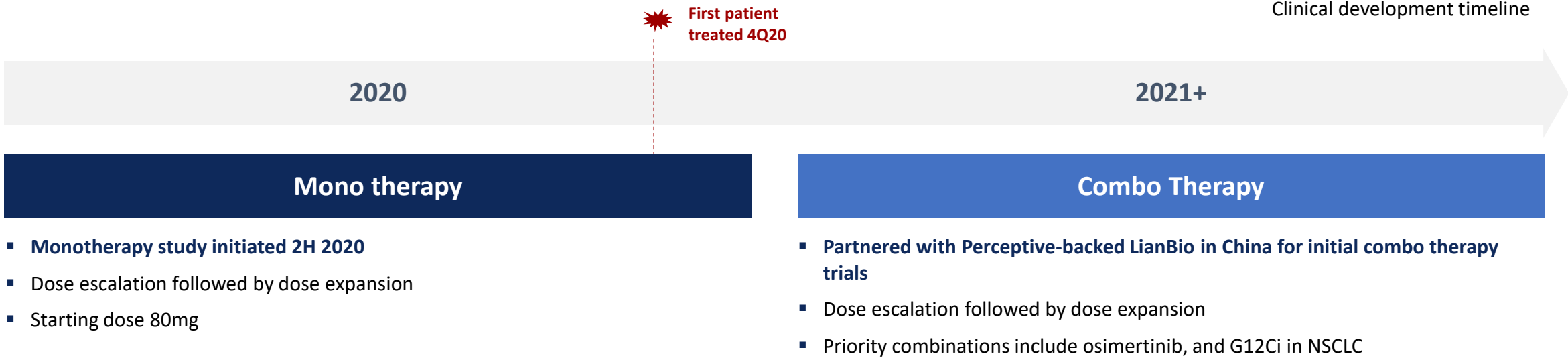
~1 million patients annually

¹ Predicted human PK based on preclinical in vivo data ² Preclinical data of combination efficacy with SHP2i

SOURCE: US incidence estimated from SEER, TCGA and Kiuru & Busam "The NF1 gene in tumor syndromes and melanoma"; all scaled for WW incidence

SHP2: BBP-398 monotherapy study initiated in 2020; combo trials to follow

Clinical development timeline



Initial clinical combinations of focus based on SHP2i preclinical data

	SHP2i Combination Partner	Tumor growth inhibition
KRAS G12Ci	AMG 510	~130%
EGFRi	Osimertinib	~125%
PD-1	Anti-mouse PD-1	~90%
MEK	Trametinib	~80%
CDK4/6 and MEK	Trametinib + palbociclib	~110%

2021 is a pivotal year with major catalysts across the pipeline

2021		ANTICIPATED
1H	2H	2022
<ul style="list-style-type: none"> ☑ BBIO / EIDX merger closure: Shareholder meeting January 19 ☑ NULIBRY (fosdenopterin) for MoCD type A: FDA approval ☐ Encaleret (CaSRi) for ADH1: Ph2 proof-of-concept data (March 20) ☐ High-dose infigratinib (FGFRi) for second-line cholangiocarcinoma: FDA approval 	<ul style="list-style-type: none"> ☐ Acoramidis (ATTR stabilizer) for ATTR-CM: Ph3 ATTRibute topline data ☐ Low-dose infigratinib (FGFRi) for achondroplasia: Ph2 proof-of-concept data ☐ AAV5 gene therapy for CAH: Initial data from first-in-human study (late '21 / early '22) ☐ COL7 replacement for RDEB: Data from Ph2 study (late '21 / early '22) 	<ul style="list-style-type: none"> ☐ Acoramidis (ATTR stabilizer) for ATTR-CM: NDA submission ☐ KRAS inhibitor program: Clinical candidate selection ☐ SHP2 inhibitor for RAS and RTK driven cancer: Monotherapy Phase 2 dose selection ☐ Ribitol for LGMD2i: Ph2 proof-of-concept data

\$607mn in cash and equivalents as of December 2020 plus ~\$750mn in gross proceed from recent convertible offering anticipated to provide runway into 2023