



## COMPANY PRESENTATION

April 2020



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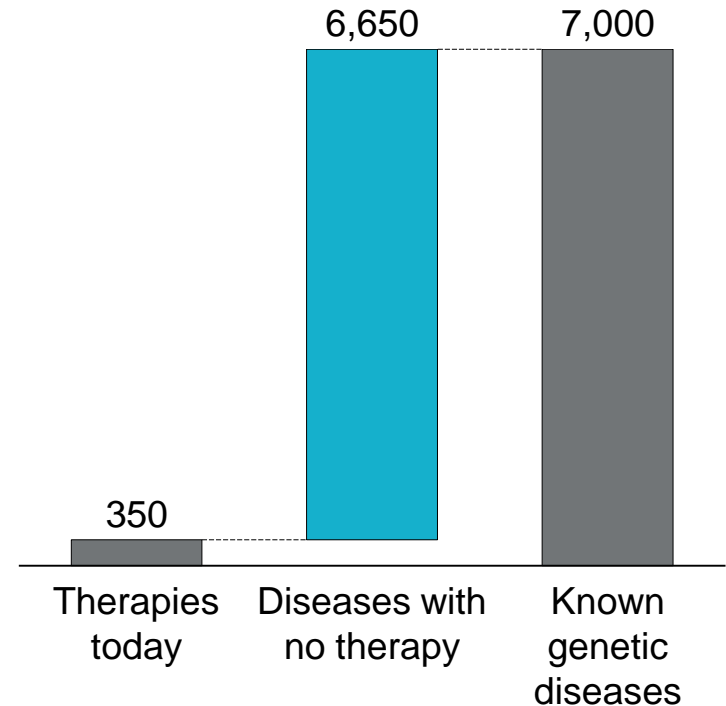
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# We are at Day 1 in the era of genetic medicine

## Advances in science and medicine (2019)

- **Better context:** Cryptic genetic variation and modifiers
- **Better understanding heterogeneity:** Genetic interaction manifolds and the wonderful story of Hirschsprung's Disease
- **Deeper saturation:** Saturation genome editing
- **Faster:** Rapid whole-genome sequencing in the ICU
- **Developing infrastructure:** National Human Genome Research Institute (NHGRI) reports cost per genome at \$942 this year (all time low)
- **Striking new therapeutics:** SCD, CF, PN, TTR, SMA, and others

## Vast opportunity to help patients



# We are building a leading genetic disease company

## Core attributes...

1. Distinctive early stage asset selection
2. Experienced, product-focused R&D team
3. Efficient corporate structure
4. The willingness and scale to fail
5. Focus at the level of individual diseases and assets

## ...applied many times...



**+ 18 BridgeBio programs**

## ...a pipeline of potential blockbusters and synthetic blockbusters\*

- Two potential \$1B+ franchises in Phase 2 or later
- Two planned NDA submissions this year
- Several early-stage potentially large franchises
  - KRAS
  - GPX4
  - Congenital adrenal hyperplasia
  - Leber's hereditary optic neuropathy
- Multiple IND submissions planned in 2020
- Four new programs announced in January 2020

\*Blockbuster defined as program with \$1bn+ opportunity

# BridgeBio is led by a world-class team of experienced drug developers

We rely on some of the top R&D minds in this industry to select assets...

**Charles Homcy, MD**  
Chairman of Pharmaceuticals



**Frank McCormick, PhD, FRS**  
Chairman of Oncology



**Richard Scheller, PhD**  
Chairman of R&D

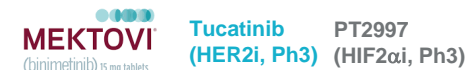


...and put them in the hands of one of the most productive groups of R&D operators in the industry

**Uma Sinha, PhD**  
Chief Scientific Officer



**Eli Wallace, PhD**  
Chief Scientific Officer in Residence, Oncology



**Robert Zamboni, PhD**  
Chemistry



Together, our R&D team is responsible for 100+ INDs and 20+ approved products

# Assessing BridgeBio

## Criteria

## Relevance

## Focus Today

1

High probability of success

- Historically higher probability of success for genetic disease drugs
- BridgeBio's early programs have outperformed historical probabilities

Current  
Pipeline  
Progress

2

Number of programs

- We find great science and unlock its potential for patients
- Always searching for the next PellePharm or Eidos
- Scale allows for objective assessment and failure

New  
Programs

3

Capital efficiency

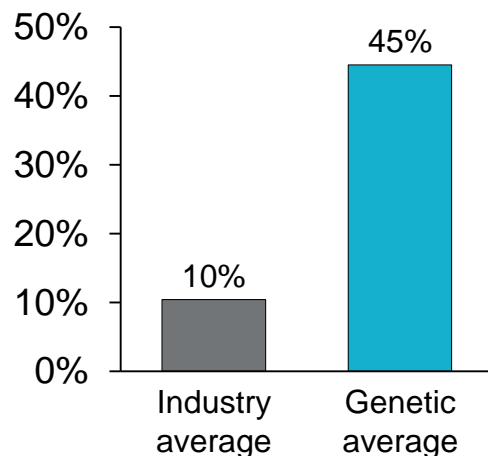
- Generate value by making programs ROI-positive
- Driven by judicious use of capital at the high-risk preclinical stages

Spend to  
IND

# We believe genetic disease drug discovery is lower risk, faster, with potentially higher returns than traditional drug discovery

**>4x** Higher cumulative probability of success

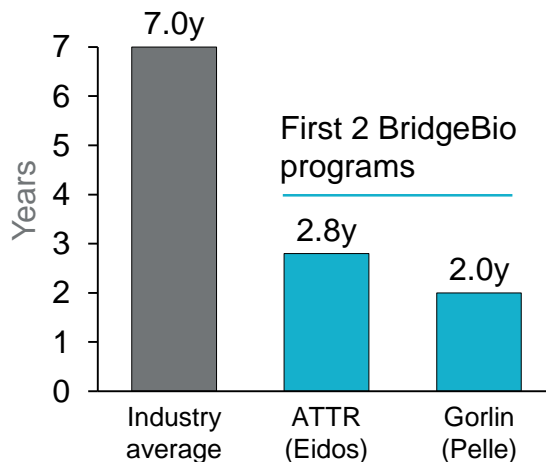
Probability of success from Ph1 to launch



Sources: Hay et al., Nature Biotechnology, "Clinical Development Success Rates for Investigational Drugs", 2014

**>65%** Faster time to Phase 3\*

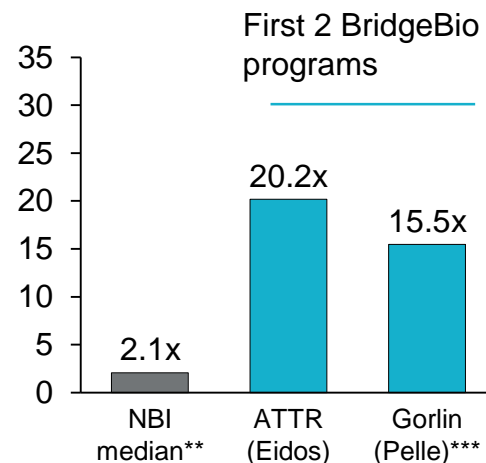
Time from lead optimization to Ph3



Sources: Paul et al., Nat Rev Drug Disc, "How to improve R&D productivity: the pharmaceutical industry's grand challenge.", 2010

**>8x** Better return on investment\*

Total return on investment  
[Enterprise value]/[APIC - cash on hand]\*



\*As of 2/28/2020 close

\*\*Includes all NBI constituents with market value <\$20bn

\*\*\*Calculated as total consideration from LEO Pharma transaction divided by total burn to date

Sources: FactSet

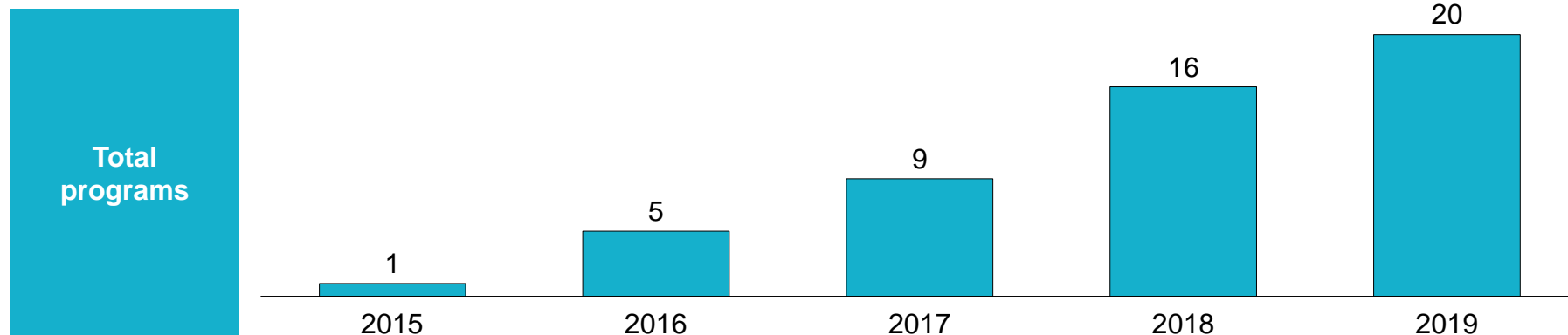
**Targeting genetic disease has higher average probability of success and BridgeBio has demonstrated higher ROI and shorter development time in its first 2 programs**

\*For first two BridgeBio programs

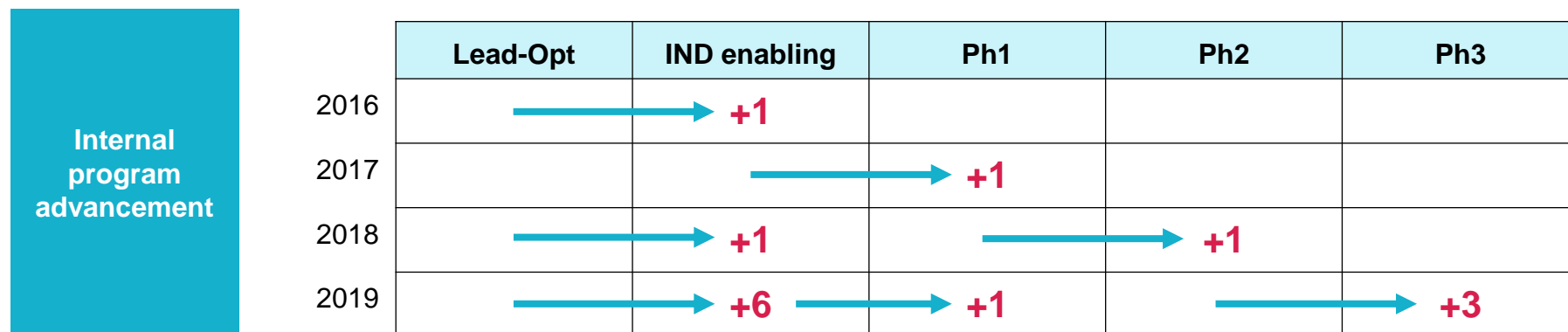
# A rapidly-advancing pipeline

Since our inception, we have actively built our pipeline through business development efforts, including the acquisition and in-licensing of assets, and advancing programs through internal stage-gates

## Growth of assets in our pipeline:



## Advancement of product candidates through key stage-gates:



# Our pipeline of 20+ development programs spans multiple therapeutic areas and drug modalities



Small molecule



Topical small molecule



Biologics



Gene therapy

Portfolio segment	Program <sup>1</sup>	Drug mechanism	Diseases	Patient pop. (US+EU) <sup>2</sup>	Modality	Pre-Clinical		Clinical		
						Discovery	IND-enabling	Phase1	Phase 2	Phase 3
<b>Mendelian</b> 	<b>AG10</b>	TTR stabilizer	ATTR-CM	>400K						
	<b>BBP-870</b>	cPMP replacement	MoCD type A	100						NDA
	<b>Infigratinib</b>	Low-dose FGFR1-3i	Achondroplasia <sup>4</sup>	55K						
	New program	<b>Encaleret</b>	CaSR antagonist	ADH1 / HP						
	New program	<b>Zuretinol</b>	Synthetic retinoid	IRD (RPE65 or LRAT)						
	<b>BBP-418</b>	Glycosylation substrate	LGMD2i	7K						
	<b>BBP-711</b>	GO1 inhibitor	PH1 / FSF	5K / 1.5M						
	<b>BBP-761</b>	Succinate prodrug	LHON	20K						
	<b>BBP-671</b>	PanK activator	PKAN / OA	7K						
	New program	<b>BBP-472</b>	PI3Kβi	PTEN autism						
<b>Genetic Dermatology</b> 	<b>Patidegib<sup>3</sup></b>	Topical SMOi	Gorlin / BCC	120K						
	<b>BBP-589</b>	Recombinant COL7	RDEB	1.5K						
	<b>BBP-681</b>	Topical PI3Kαi	VM / LM	117K						
	<b>BBP-561</b>	Topical KLK 5/7i	Netherton	11K						
<b>Targeted Oncology</b> 	<b>Infigratinib</b>	FGFR1-3i	FGFR+ tumors	37K						
	<b>BBP-398</b>	SHP2i	Multiple tumors	>500K						
	<b>BBP-454</b>	Pan-mutant KRASi	KRAS+ tumors	>500K						
	<b>BBP-954</b>	GPX4i	Multiple tumors	>500K						
<b>Gene Therapy</b> 	<b>BBP-631</b>	21-OH gene therapy	CAH	>75K						
	<b>BBP-812</b>	ASPA gene therapy	Canavan	1K						
	New program	<b>BBP-815</b>	TMC1 gene therapy	Genetic hearing loss						

<sup>1</sup> Each of our programs is housed in a separate subsidiary; <sup>2</sup> Patient population: Prevalence except for asterisked figures which represent incidence; <sup>3</sup>We are party to an option agreement pursuant to which LEO Pharma A/S has been granted an exclusive, irrevocable option to acquire PellePharma, 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. <sup>4</sup>Protocol accepted by Australian local ethics committee, IND submission to FDA expected 2020.

# 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 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

Claudia, child with achondroplasia

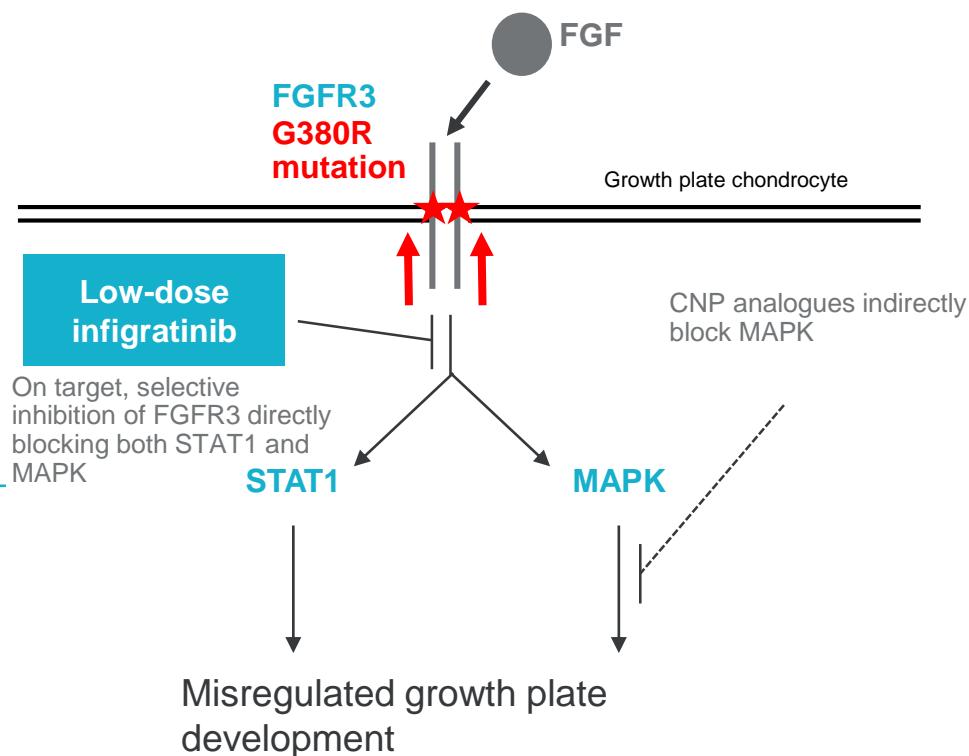
# Potential best-in-class approach to treating achondroplasia directly at its genetic source

## 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 causes cranial, spinal, and stature symptoms

## Low-dose infigratinib has the potential to:

- Directly inhibit the causal gain-of-function mutation in FGFR3
- Normalize both the STAT1 and MAPK signaling pathways
- Demonstrate clear macro and microscopic improvements on foramen magnum, intervertebral discs, and long bones in validated preclinical model



# 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

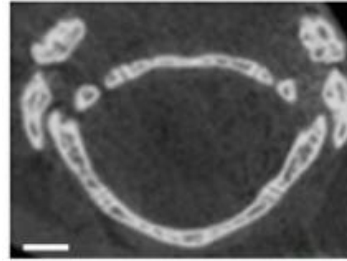
**6%**

increase in AP  
skull length

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

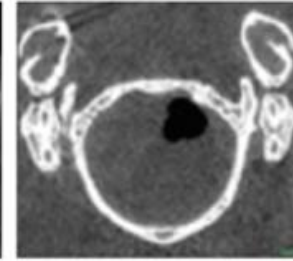
**FGFR3 WT**

No treatment



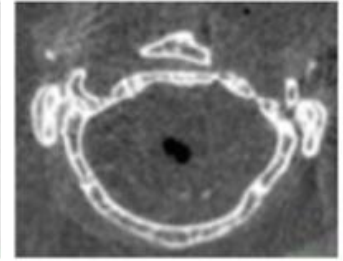
**FGFR3<sup>Y367C/+</sup>**

No treatment



**FGFR3<sup>Y367C/+</sup>**

**Infigratinib tx**



## 2 Disorders of the spine

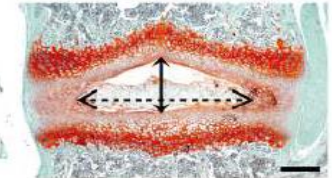
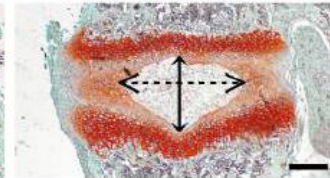
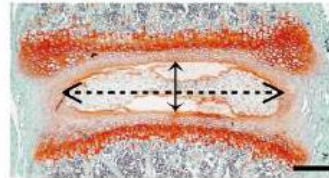
**12%**

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

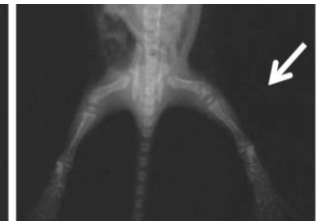
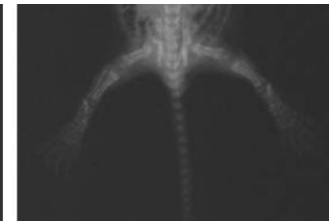
**21%**

increase in  
femur length

**33%**

increase in  
tibia length

May lead to **increased stature** and **proportionality**



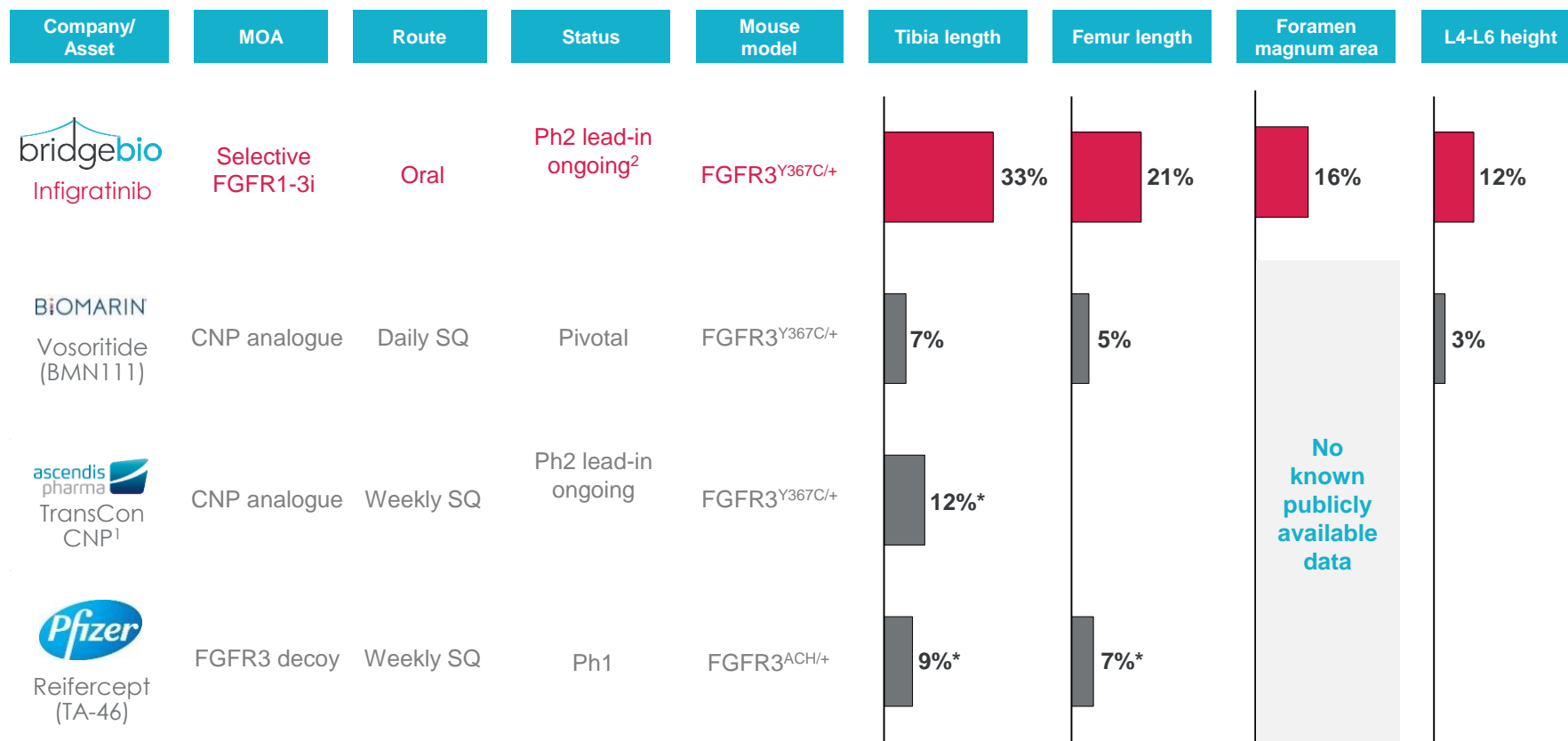
Source: Komla-Ebri et al. J Clin Inv 2016

Note: percent increase compared to vehicle treated FGFR3<sup>Y367C/+</sup> mouse, infigratinib treatment with 2mg/kg subcutaneous dose

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

## Preclinical data from infigratinib and other investigational achondroplasia therapies

Percent increase compared to non-treated mouse



Source: Komla-Ebri et al. J Clin Invest 2016, Lorget et al. Am J Hum Genet 2012, Garcia et al. Science Trans Med 2013, Breinholt ENDO 2017

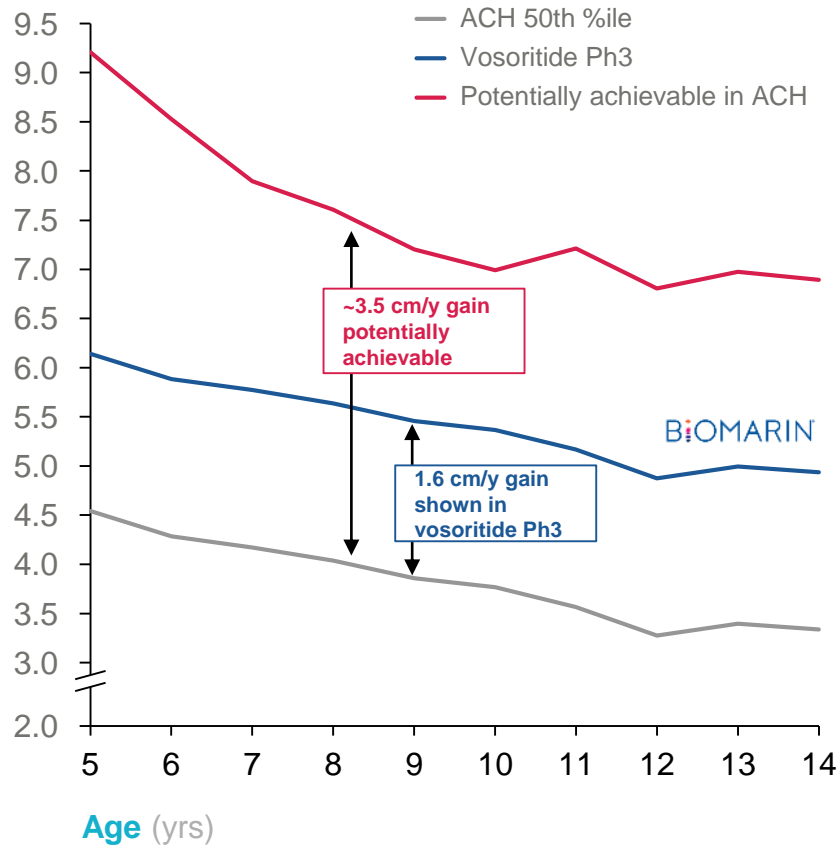
Note: subcutaneous doses, percent increase compared to vehicle treated FGFR3<sup>Y367C/+</sup>, FGFR3<sup>ACH/+</sup> mouse as noted in "Mouse model" columns  
Infigratinib treatment with 2mg/kg subcutaneous dose

<sup>1</sup>Based on vosoritide continuous infusion; \*Value estimated using Digitizelt. <sup>2</sup>Protocol submitted to Australian local ethics committee, IND submission to FDA expected 2020.

# Efficacy of CNP analogues observed to date still allows for improvements with new treatments

## Annualized growth velocity

Cm/yr



## End adult height in feet<sup>1,2</sup>

Projected height using AGV

**5'4"** Potential end adult height with treatment that increases AGV to 95<sup>th</sup> %ile of average stature from ages 5-14

**4'7"** Projected end adult height for children treated with vosoritide from ages 5-14 without tachyphylaxis

**4'1"** Average end height for people with achondroplasia who do not undergo treatment (e.g., limb lengthening)

<sup>1</sup> End adult height defined as projected 14 year old height; <sup>2</sup> Average height across genders

NOTE: vosoritide AGV shown reflects a mean increase over baseline of 1.6 cm/yr, interpreted as a mean increase of 1.6 cm/yr over 50th percentile at every age;

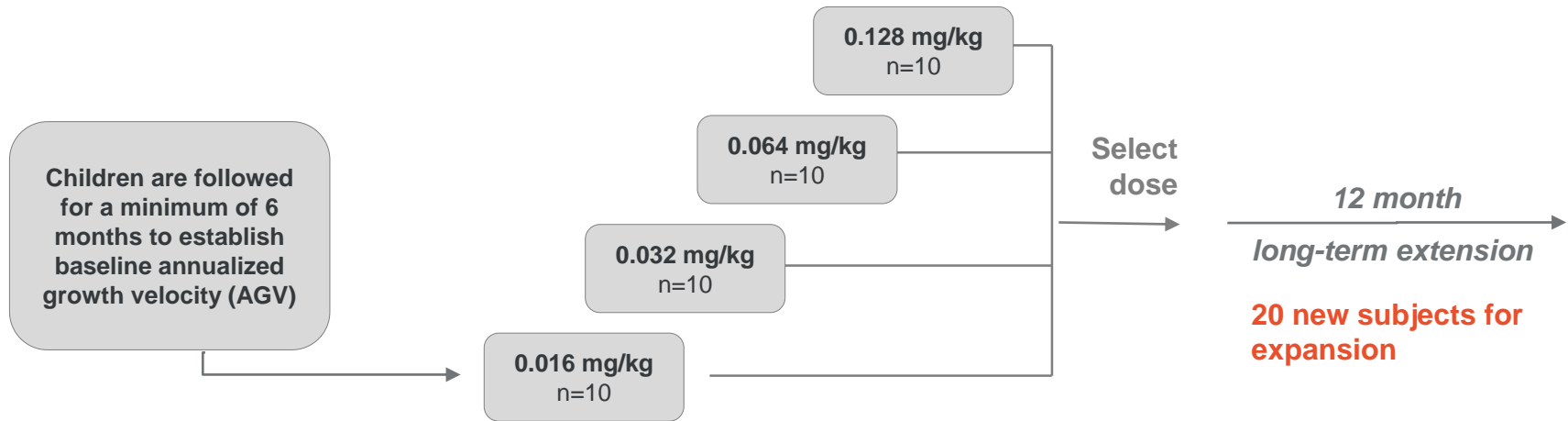
SOURCE: Hoover-Fong et al 2017, Hoover-Fong et al 2008, Savarirayan et al NEJM 2019

# The PROPEL clinical program is enrolling and potential POC data expected in 2021

Observational run-in

Ph2 Dose-finding (n=40)

Expansion (n=20)



## 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

# Recombinant collagen type VII for recessive dystrophic epidermolysis bullosa (RDEB)



## RDEB overview:

- **Prevalence:** 1,500 (US + EU)
- **Genetic driver:** mutations in the COL7A1 gene encoding the protein collagen type VII
- **Pathophysiology:** Systemic impairment of dermal-epithelial cohesion throughout various tissues leading to painful blistering on the skin, GI tract, and oral cavity

## Features of a potential best-in-class medicine for RDEB:

- **Targeting RDEB at its genetic source**, by replacing missing COL7 protein via a simple IV infusion
- **Potential to address burden of RDEB beyond the skin**, including systemic manifestations
- **Proactively address wound formation and healing**, rather than reactively treat lesions

Bardy, child with RDEB

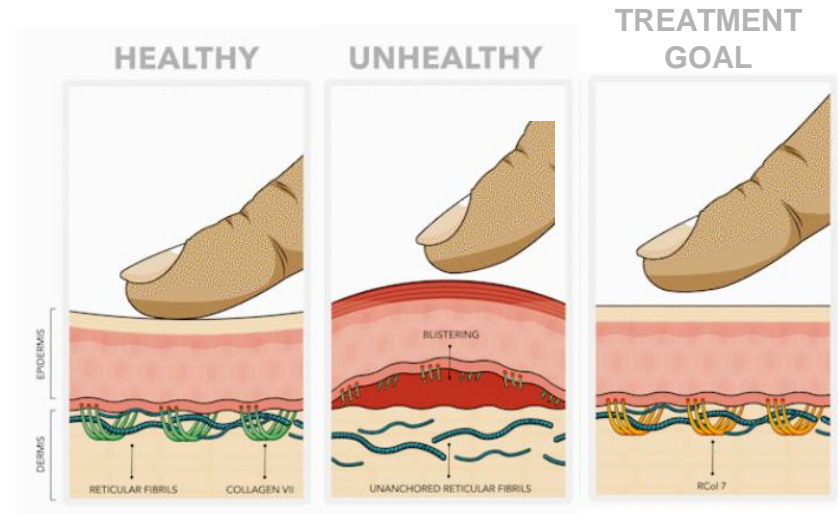
# Recombinant collagen type VII for recessive dystrophic epidermolysis bullosa (RDEB)

## RDEB COL7 loss-of-function mutations cause:

- Near complete loss of COL7 at epithelial junctions on the skin and throughout the body
- Painful erosions and blistering on the skin, GI tract, and oral cavity
- Failure to thrive, decreased life span, high risk for squamous cell carcinoma

## Our systemic COL7 replacement is designed to:

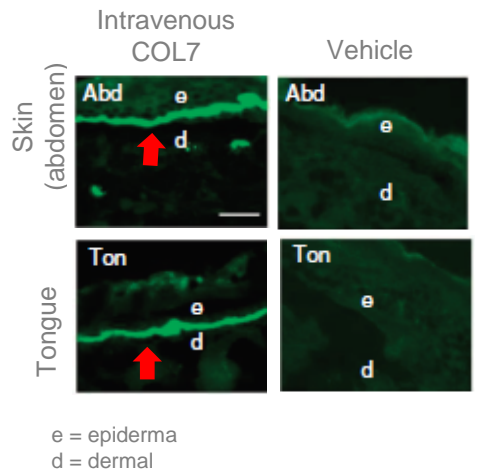
- Replace COL7 at epithelial junctions throughout the body
- Address the systemic burden of RDEB including on the skin, GI tract and oral cavity
- Proactively address wound formation and healing globally rather than reactively treat lesions



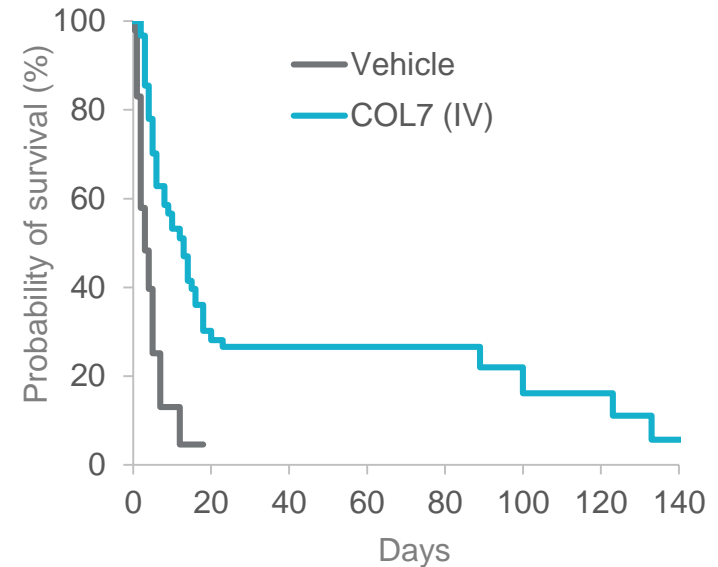
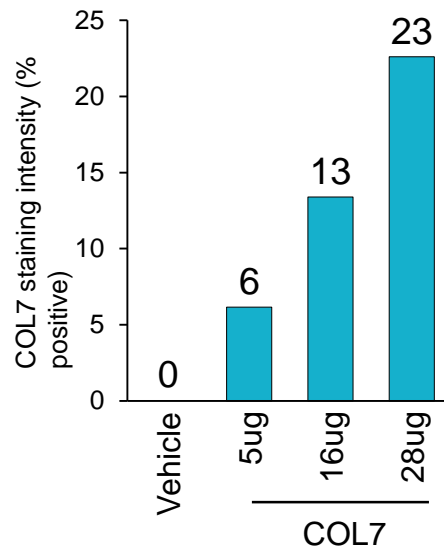
# Recombinant COL7 distributes systemically, leading to survival benefits in the RDEB mouse model

A single intravenous injection of recombinant COL7 distributed to epithelial barriers throughout the body (skin, oral cavity, GI tract), in a dose-dependent manner

This led to a **significant survival benefit** in COL7-treated animals



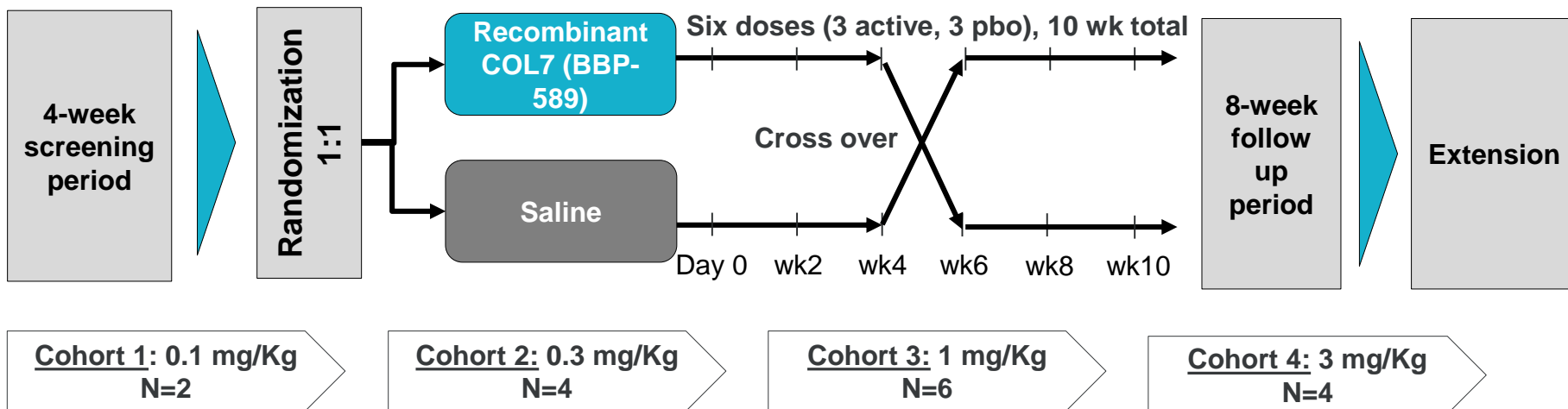
COL7 is stained green in images above



# Ongoing randomized, dose-escalation Phase 1/2 proof-of-concept clinical study in adults with RDEB

- First patient dosed in 1Q19
- [Anticipate potential POC data in 2020](#)

## Protocol for EACH cohort



## KEY INCLUSION CRITERIA

- Adult with RDEB diagnosis
- Deficiency but not total loss of COL7 protein
- At least 1 wound >20cm<sup>2</sup> for ≥6 weeks

## KEY EXCLUSION CRITERIA

- Known hypersensitivity to BBP-589
- Received investigational RDEB agent in last 6 months

## PRIMARY ENDPOINT

- Safety and tolerability

## KEY SECONDARY AND EXPLORATORY ENDPOINTS

- COL7 deposition and residence time in skin biopsies
- Change in healing of up to 5 target wounds
- Patient reported outcomes (itch, QoL)

# Targeted oncology portfolio



Andrea,  
CCA patient

## World-class oncology team drives our discovery and development

- **Eli Wallace:** CSO Oncology
- **Frank McCormick,** Chairman of Oncology
- **Richard Scheller,** Chairman of R&D



Genentech



## We target driver mutations in genetically defined cancers...

- **FGFR1-3i** for FGFR+ cancer: Near-term revenue in CCA, multiple expansion indications
- Pan-mutant **KRASi** for KRAS+ cancer: Platform approach in partnership with NCI RAS initiative

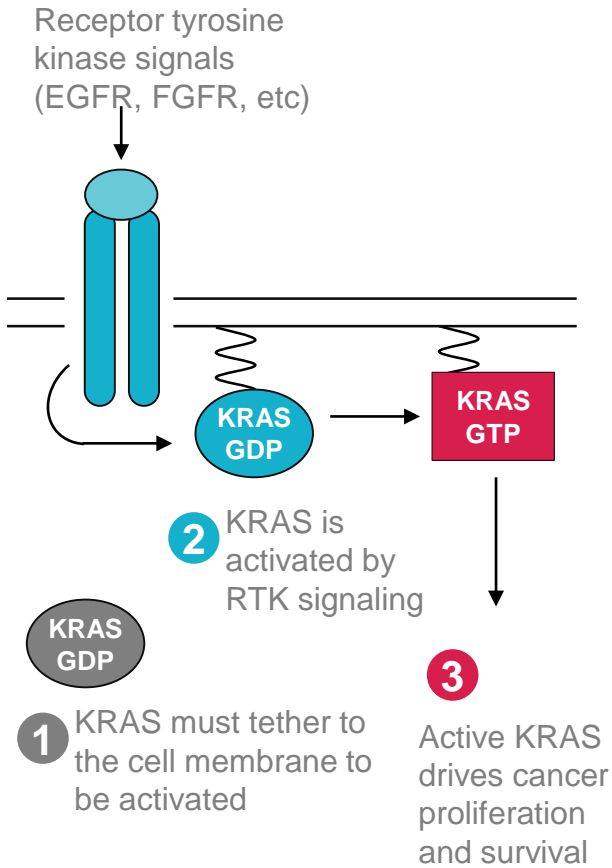
## ...while also focusing on novel targets with extensive academic validation

- **SHP2i** for multiple tumors (10+ recent papers in *Nature*, *Science*, *Nature Medicine*)
- **GPX4i** for multiple tumors (10+ recent papers in *Nature*, *Cell*, *Cancer Cell*)

Program	MOA	Disease	Stage	Next anticipated update
Infigratinib	FGFR1-3 inhibitor	FGFR+ cancer	Ph3	Pivotal CCA data 2020, NDA 2020
BBP-398	SHP2 inhibitor	Multiple tumor types	Pre-IND	IND submission in 2020
BBP-454	Pan-mutant KRAS inhibitor	KRAS+ cancer	Discovery	Clinical candidate nomination
BBP-954	GPX4 inhibitor	Multiple tumor types	Discovery	Clinical candidate nomination

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

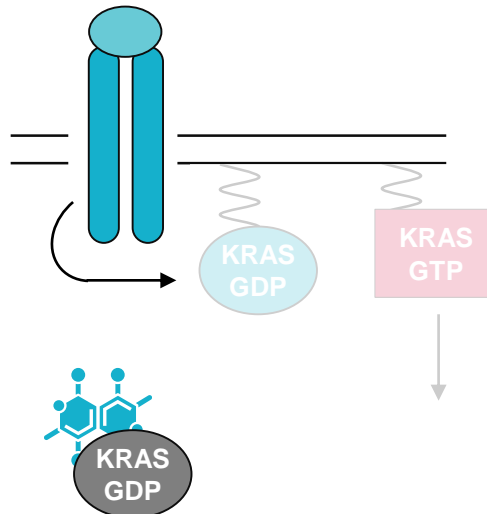
## KRAS activation in cancer is a multi-step process



## Our programs target different steps of the KRAS activation process

### Program 1: C185 targeting

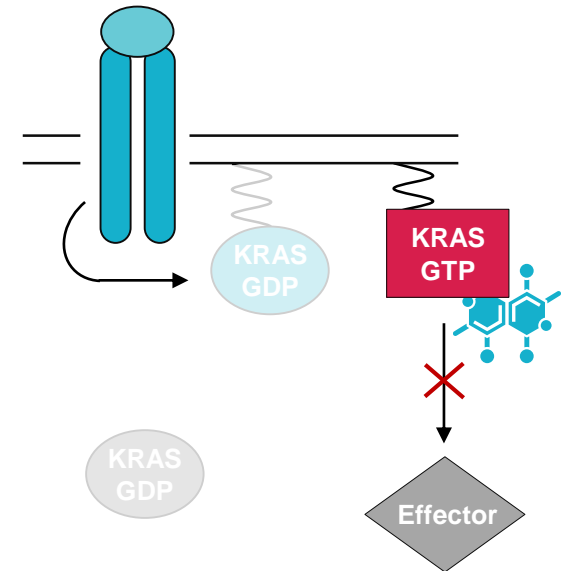
- Blocks KRAS from tethering
- Blocks conversion of inactive KRAS GDP to active KRAS GTP



KRAS tethering is blocked – cancer growth is inhibited

### Program 2: H95 targeting

- Directly binds activated KRAS
- Inhibits KRAS from signaling through effectors



Activated KRAS signaling is inhibited

# SHP2: Our potentially best-in-class SHP2 inhibitor is expected to enter the clinic mid-2020

- SHP2 connects RTK signaling to downstream MAPK signaling activation
- Our compound potently traps SHP2 in an inactive state, thereby potentially blocking downstream oncogenic signaling
- In collaboration with MD Anderson, optimized our SHP2i for use in combination and reduced cardiac liability
  - No evidence of QTc prolongation or hypertension
- BBP-398 was well tolerated in rats and dogs in 28d GLP-tox studies
  - Histological and clinical chemistry findings consistent with MAPKi
  - At maximum doses (25 mg/kg/day, dogs; 100 mg/kg/day, rats), MTD was not reached
- IP published 02/13/2020
- First SHP2 inhibitor clinical data, (RVMD Q1 2020) demonstrates monotherapy antitumor activity\*

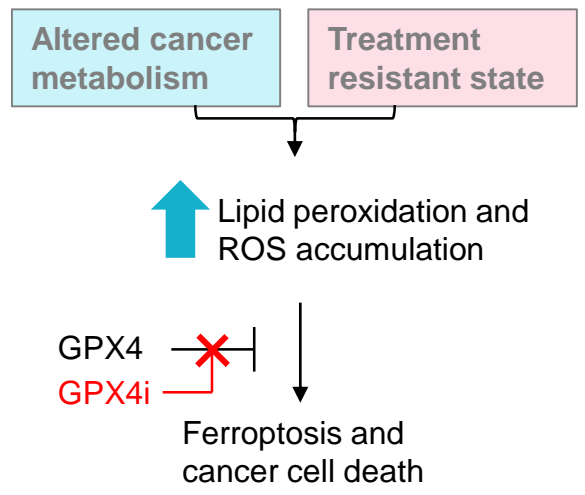
## Preclinical SHP2i data

<i>In vitro</i> properties	BBIO	RVMD*
pERK IC <sub>50</sub> (nM) cellular assay	<40	<40
hERG Patch clamp IC <sub>50</sub> (μM)	>100	?
Monotherapy anti-tumor activity		
KRASG12C xenograft	✓	✓
EGFR mutant xenograft	✓	✓
Combination enhanced anti-tumor activity		
G12Ci	✓ (AMG 510)	✓ (MRTX 849)
MEKi	✓ (trametinib)	✓ (cobimetinib)
EGFRi osimertinib	✓	✓

**Preclinical profile demonstrates activity in-line with SHP2i class and potential for better tolerability**

# GPX4: Potential first-in-class therapy for a novel cancer target

## MOA

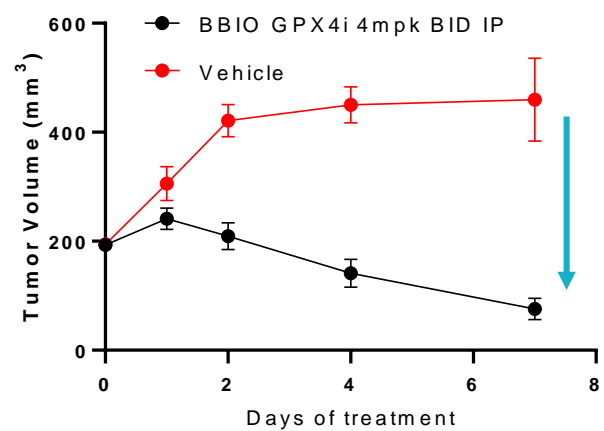


- GPX4 was recently identified as a key tumor dependency in multiple CRISPR screens and mechanistic studies
- GPX4 allows tumor cells to survive by neutralizing toxic lipid peroxides
- Our approach is to directly inhibit GPX4, thereby triggering cancer death through ferroptosis

## Key data

### In vivo monotherapy activity in a renal cell carcinoma mouse model

Model: 786-O RCC xenograft (VHL LOF, p53 LOF)



Rapid tumor regression after only 7 days of dosing

### Synergy with targeted therapies and immunotherapy using in vitro models

# Infigratinib (FGFRi): Near-term submission in CCA and multiple large expansion opportunities

Indication	Key Data	Status	Next planned update
<b>FGFR2+ cholangiocarcinoma</b>	<b>39% ORR</b> in patients with $\leq 1$ previous line of treatment	<ul style="list-style-type: none"><li>Enrollment complete in 2L Ph2 pivotal cohort</li><li>Ph3 in 1L study enrolling</li></ul>	<ul style="list-style-type: none"><li>Updated pivotal data 2H20</li><li>NDA submission 2H20</li><li>2021 launch</li></ul>
<b>FGFR3+ urothelial carcinoma</b>	<b>25% ORR</b> in metastatic relapsed refractory setting suggests clear activity in this tumor	<ul style="list-style-type: none"><li>FPI for Ph3 in adjuvant setting in 1H20</li></ul>	<ul style="list-style-type: none"><li>Complete enrollment in Ph3 adjuvant study</li></ul>
<b>FGFR fusion-positive tumor agnostic</b>	<b>5 tumor types</b> Showed response to infigratinib in Ph1/2	<ul style="list-style-type: none"><li>FPI for Ph2 signal optimization study in 1H20</li></ul>	<ul style="list-style-type: none"><li>Potential Ph2 data 2021</li></ul>

# Gene therapy portfolio



Vayle, child with Canavan

## Experienced team with track record in gene therapy



BIOMARIN



NOVARTIS

## Partnered with top academics in the gene therapy space

- Guangping Gao, Ph.D (UMass)
- Pierre Bougneres, M.D., Ph.D. (INSERM)
- Jeff Holt, Ph.D (Boston Children's)

## Congenital adrenal hyperplasia (BBP-631)

- One of the largest known AAV gene therapy markets (prevalence 75K US+EU)
- Low threshold to correct phenotype, validated by human genetics
- Durable transgene delivery and expression for 6-month in NHP study

## Canavan disease (BBP-812)

- Lethal, degenerative, neuromuscular disease
- Precedented AAV9 serotype with safety data in one compassionate use case

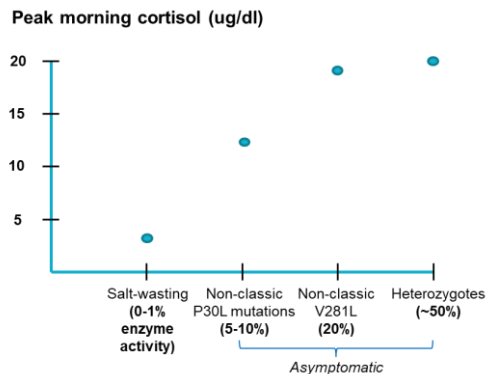
## TMC1-driven hearing loss (BBP-815)

- Delivers functional copy of TMC1 gene allowing transmission of auditory stimuli
- Nature Communications publication shows significant rescue of hearing function in diseased mice

Program	MOA	Disease	Stage	Next anticipated update
BBP-631	21-OHase gene therapy	Congenital adrenal hyperplasia	Pre-IND	IND submission in 2020
BBP-812	ASPA gene therapy	Canavan disease	Pre-IND	IND submission in 2020
BBP-815	TMC1 gene therapy	Genetic hearing loss	Discovery	Clinical candidate nomination

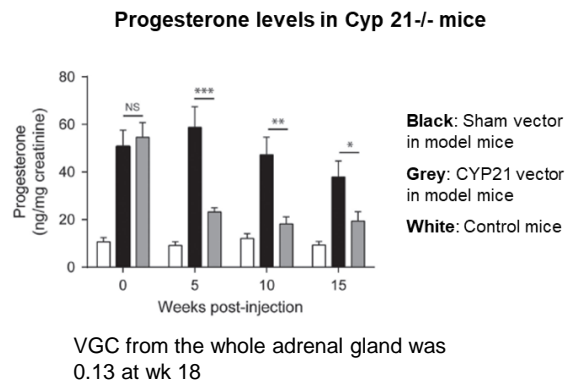
# 21-OH gene therapy for CAH: NHP study showed durable transgene expression; 5-10% of WT enzyme may be sufficient for clinical impact

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



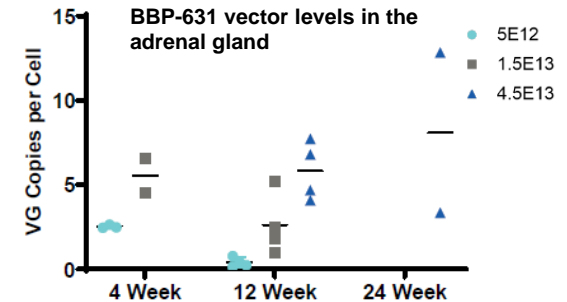
- Due to the high enzymatic efficiency/selectivity of 21-OHase, **only a small amount of enzyme is required to rescue the phenotype**

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

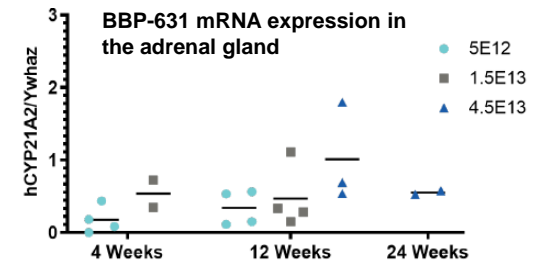


- At 15 weeks in treated mice, **progesterone** (the key substrate of 21OHase in mice) was **significantly reduced vs untreated mice**

## NHP studies show sustained VGC and RNA out to 6 months



- Mean vector genome copies per cell appear stable at 24 wks

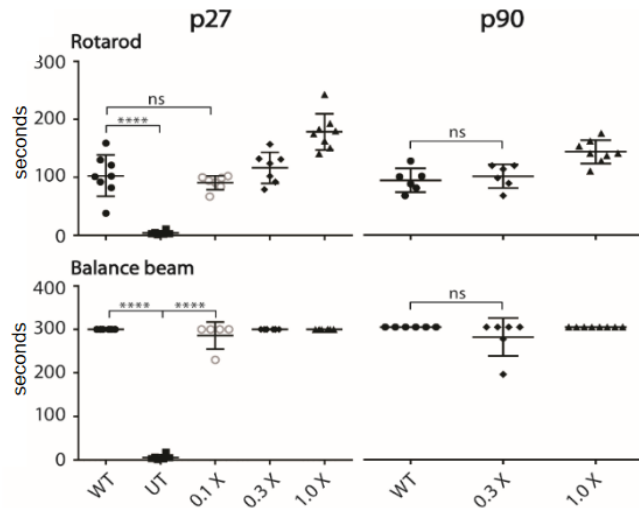


- Transgene expression is dose-dependent and stable out at 24 wks

# ASPA gene therapy for Canavan: Phenotypic correction in a lethal mouse model and broad CNS transduction in NHPs

## Mouse studies show that BBP-812 can achieve phenotypic reversal

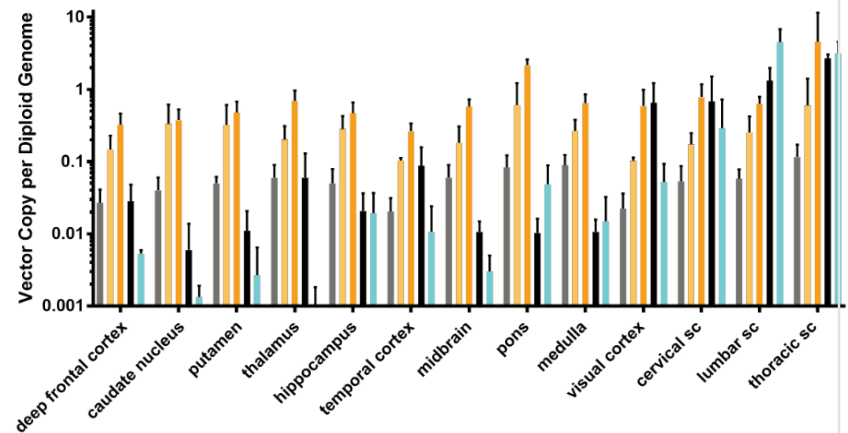
### Effect of BBP-812 on rotarod and balance beam, ASPA KO mice (untreated vs 3 different doses) and WT mice



- ASPA KO mice treated with at least  $2.6 \times 10^{13}$  vg/kg had NAA metabolism and performance on motor function tests restored. Mice treated at  $2.6 \times 10^{14}$  vg/kg outperformed WT mice.

## NHP studies show broad CNS distribution with IV delivery

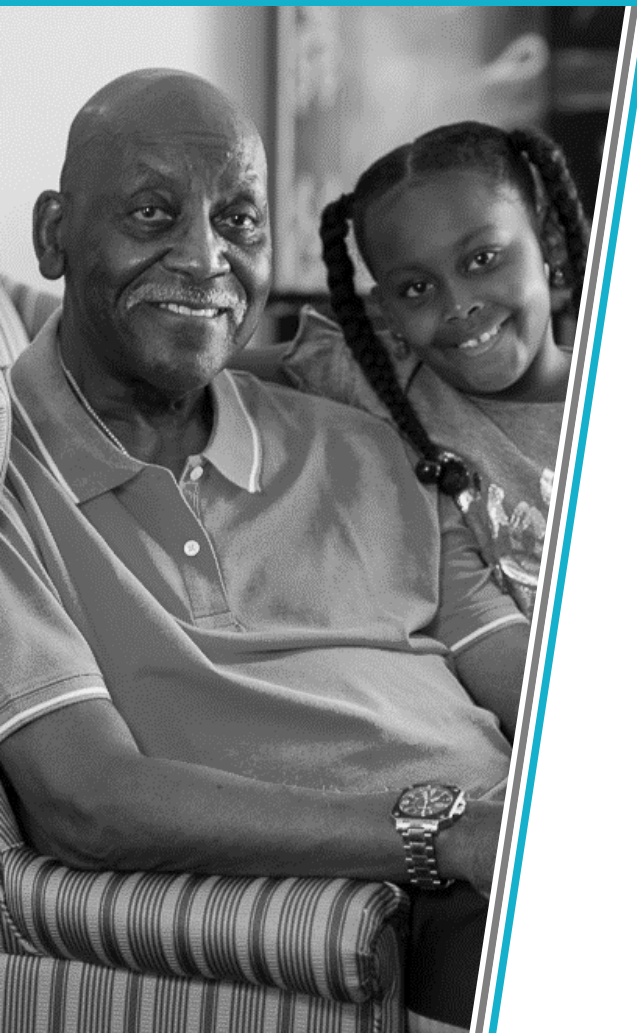
### Vector copies per diploid genome in various CNS regions in NHPs



- IV delivery of BBP-812 showed superior transduction of several CNS regions compared to ICV and IT delivery

IV-Low  
IV-Mid  
IV-High  
ICV  
IT

# AG10 for TTR amyloidosis (Eidos)

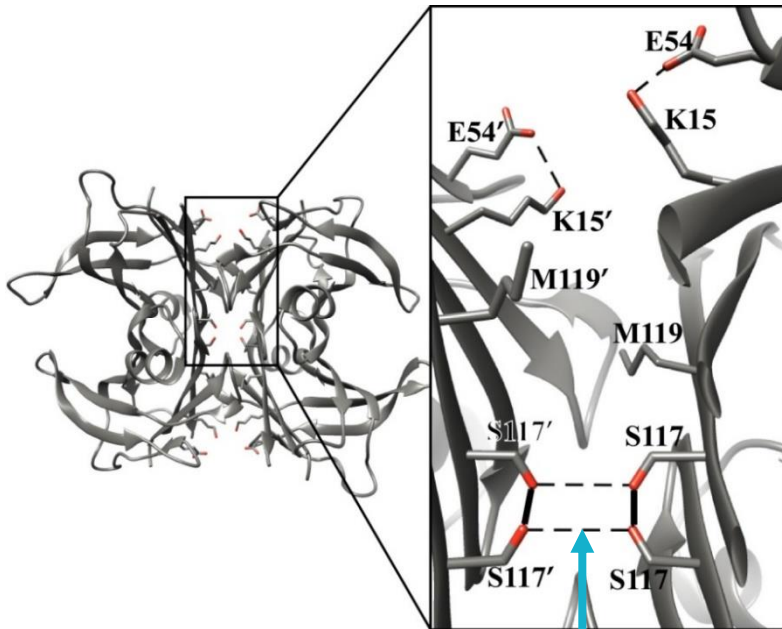


- **Addressing large and growing need in ATTR**, a fatal disease affecting >400K patients
- **Targeting the disease at its source** by stabilizing TTR, a genetic and clinically validated mechanism
- **Advancing AG10, a potential best-in-class drug** that mimics naturally occurring rescue mutation
- Phase 2 open label extension study suggests potential to **reduce mortality and cardiovascular hospitalizations** at 15 months
- **Executing Phase 3 study in ATTR-CM** with top-line data expected in 2021

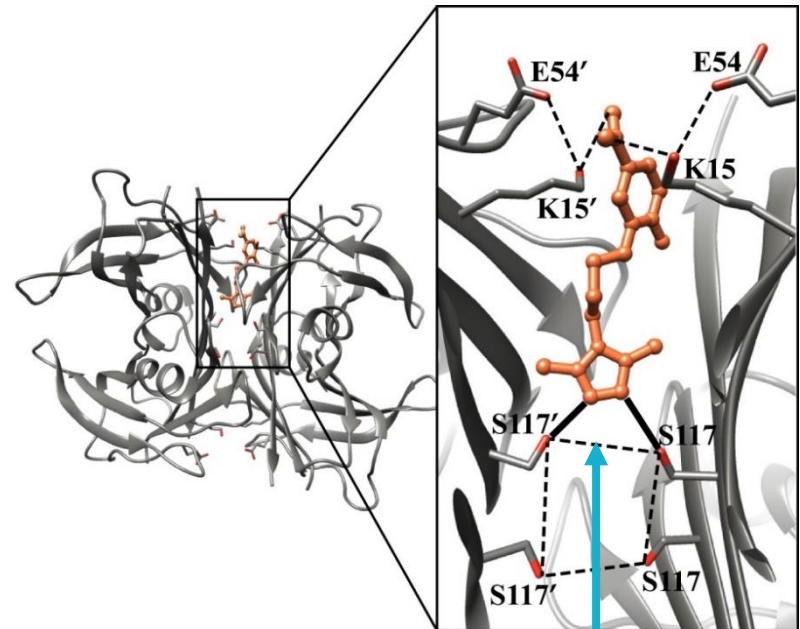
Art, ATTR-CM patient

# AG10 structurally mimics disease-protective mutation by hyper-stabilizing TTR

Disease-protective T119M mutation



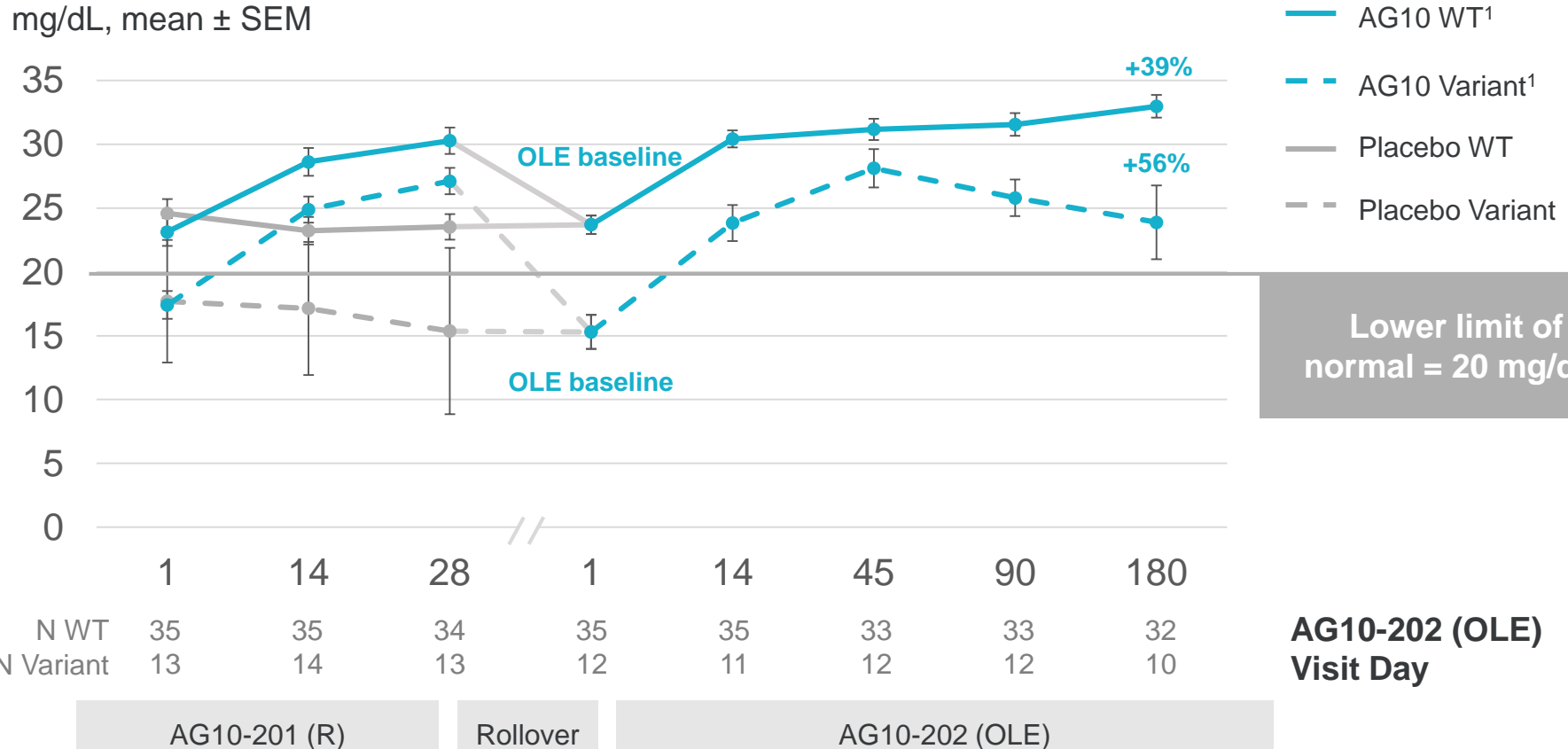
AG10-bound TTR



**Strong inter-monomer H-bonds observed via X-ray crystallography**  
**Unique binding mode vs other stabilizers**

# Serum TTR levels, a prognostic indicator of survival, increased upon AG10 treatment and were maintained throughout Ph 2 study

**Serum TTR concentration**  
mg/dL, mean  $\pm$  SEM



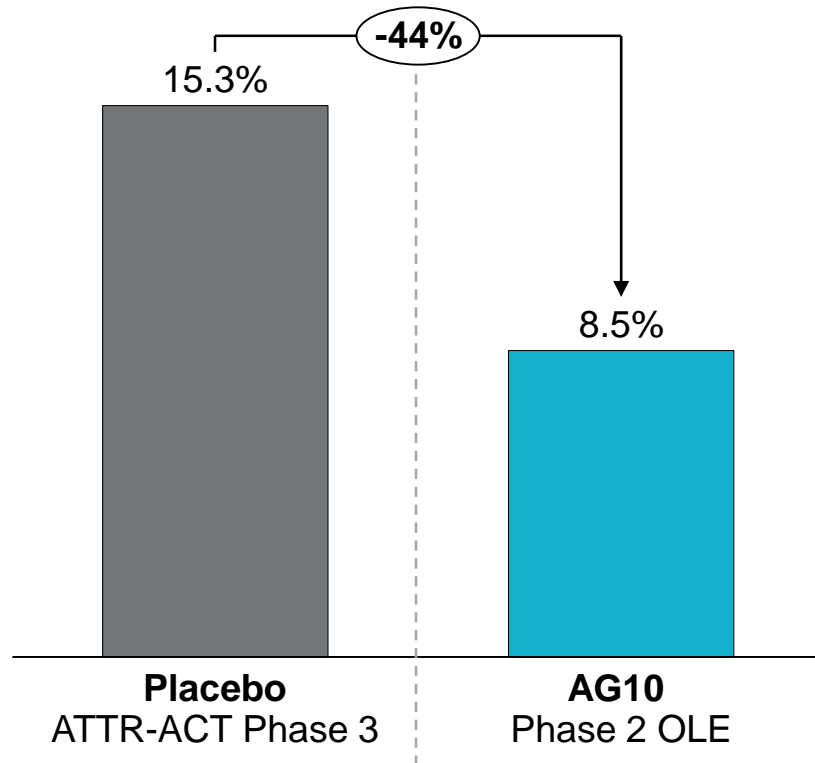
<sup>1</sup> 400mg and 800mg BID AG10 groups pooled during randomized portion

<sup>2</sup> Defined as the lower limit of the reference interval for the serum prealbumin (TTR) clinical laboratory assay

# Deaths and CV hospitalizations reported in AG10 Phase 2 OLE were lower than in placebo-treated ATTR-ACT participants

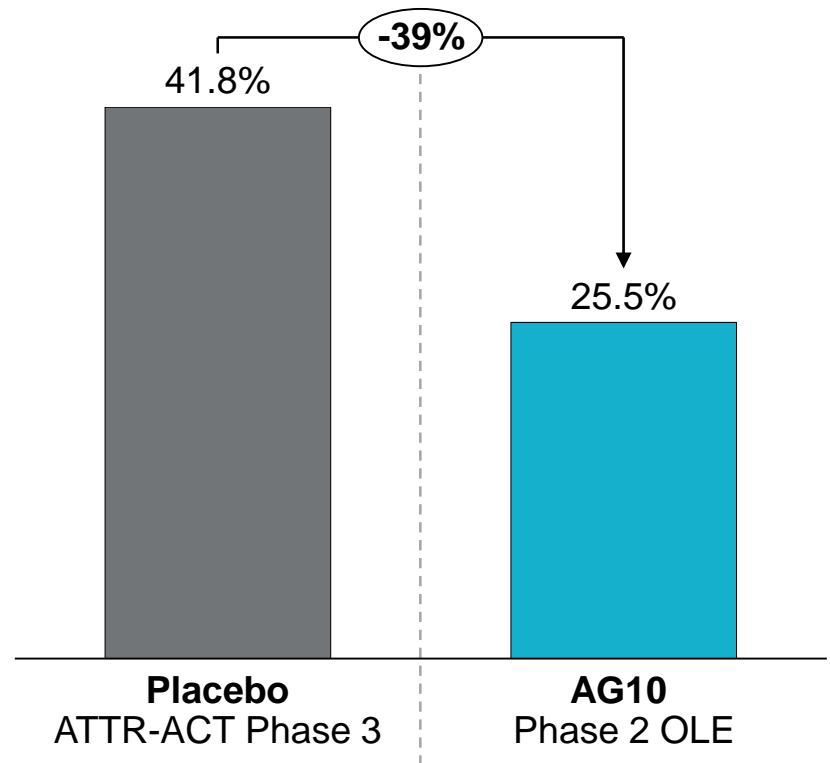
## All-cause mortality at 15 months

Participants died or receiving transplant (%)



## Cardiovascular hospitalizations at 15 months

Participants with ≥1 CV hospitalization (%)



**Phase 3 ATTRibute study expected to complete enrollment in 2H20**

<sup>1</sup> 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

# Fosdenopterin (cPMP replacement) for MoCD type A



Genetic driver: MOCS1 / cPMP depletion  
Prevalence (US + EU): 100

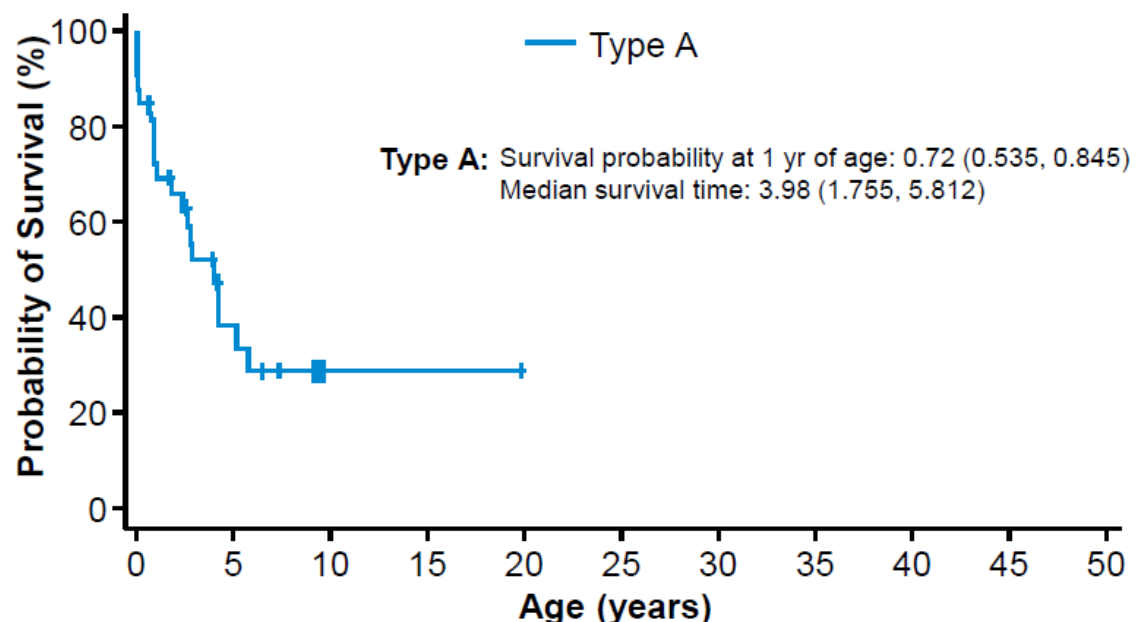
- **Designed to address an extreme unmet medical need in molybdenum cofactor deficiency (MoCD) type A**, a progressive and rapidly fatal CNS disorder (median survival < 4 years)
- **Targeting the disease at its source** by directly replacing cPMP, the missing metabolite that causes CNS toxicity
- **Potentially life saving investigational drug** with compelling pivotal data showing prolonged survival, seizure control and ambulation vs natural history
- **Rolling NDA submission initiated in 4Q19**, under FDA Breakthrough Therapy Designation

Elliott, child with  
MoCD type A

# We presented data from our natural history study in MoCD type A at SSIEM 2019

- Median survival time of <4 years highlights urgent need for a new medicine
- Data will play an important role in our NDA data package

## Kaplan-Meier Estimates of Survival Probability (FAS)



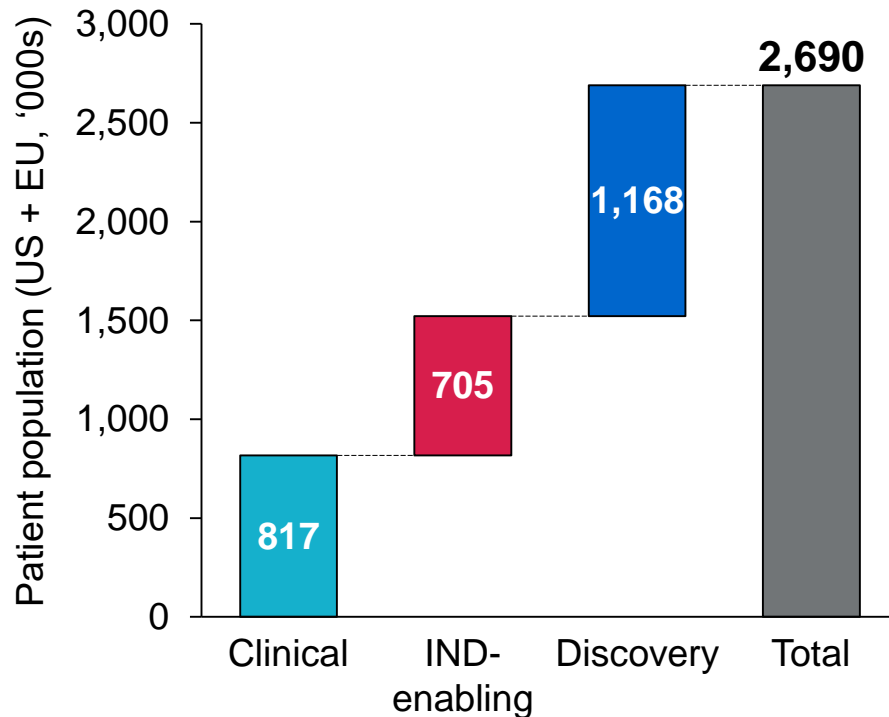
### No. patients/Patients at risk

Type A: 0/33 18/8 20/1 20/1 20/0 20/0 20/0 20/0 20/0 20/0 20/0

Note: Kaplan-Meier curves step down at time points at which a death has been observed; slashes represent patients whose observation time was censored as of the last contact date.

# Our current pipeline has the potential to treat nearly 3 million patients in the US and EU alone

## Patient population by development stage



## Breakdown of clinical-stage assets

Indication	Population
ATTR	400,000
Hypoparathyroidism	200,000
Basal cell carcinoma	120,000
Achondroplasia	55,000
FGFR+ cancer	37,000
Inherited retinal dystrophy	3,000
RDEB	1,500
MoCD type A	100

**Total:** 817,000

Our product platform has the potential to deliver diversified and sustainable revenue growth beginning in 2021

# Assessing BridgeBio

## Criteria

## Relevance

## Focus Today

1

High probability of success

- Historically higher probability of success for genetic disease drugs
- BridgeBio's early programs have outperformed historical probabilities

Current  
Pipeline  
Progress

2

Number of programs

- We find great science and unlock its potential for patients
- Always searching for the next PellePharm or Eidos
- Scale allows for objective assessment and failure

New  
Programs

3

Capital efficiency

- Generate value by making each program ROI-positive
- Driven by judicious use of capital at the high-risk preclinical stages

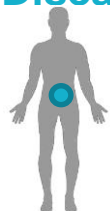
Spend to  
IND

# We recently announced four new programs including two entering Phase 2 trials

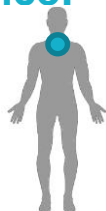
## Encaleret

**Mechanism:** Ca sensing receptor antagonist

**Diseases and prevalence:**



Autosomal dominant  
hypocalcemia type 1  
2,000  
US + EU



Hypoparathyroidism  
200,000  
US + EU

**Modality:** Small molecule



Phase 2 ready

## Zuretinol

**Mechanism:** Synthetic retinoid

**Diseases and prevalence:**



Inherited retinal disease  
caused by RPE65 or LRAT  
mutations  
3,000  
US + EU

**Modality:** Small molecule

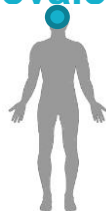


Phase 2 ready

## BBP-472

**Mechanism:** PI3K $\beta$  inhibitor

**Diseases and prevalence:**



*PTEN* autism  
120,000  
US + EU

**Modality:** Small molecule



Discovery

## BBP-815

**Mechanism:** TMC1 gene therapy

**Diseases and prevalence:**



Genetic hearing loss  
10,000  
US + EU

**Modality:** Gene therapy



Discovery

We plan to announce multiple additional new programs in 2020

# Encaleret (CaSR antagonist) for hypoparathyroidism



## Encaleret targets disease at its source by selectively antagonizing the CaSR, a key regulator of calcium homeostasis

- Opportunity to develop encaleret was identified in collaboration with global experts at the NIH
- Being prosecuted by the BridgeBio cardiorenal group



## Encaleret is a potential 1st in class CaSR antagonist with differentiated profile for hypoparathyroidism

- Initial genetically-defined population of autosomal dominant hypocalcemia type 1 (ADH1), provides high probability of success
- Potential for expansion into broader hypoparathyroidism indication (~200K patients in US & EU)



## Prior clinical experience with encaleret enables accelerated clinical development

- Well tolerated in >1,200 human subjects and increased serum calcium in a dose-dependent manner
- IND application submitted in late 2019, currently Phase 2 ready
- Proof-of-concept data in ADH1 expected in 2021

# Assessing BridgeBio

## Criteria

## Relevance

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Current Pipeline Progress

2

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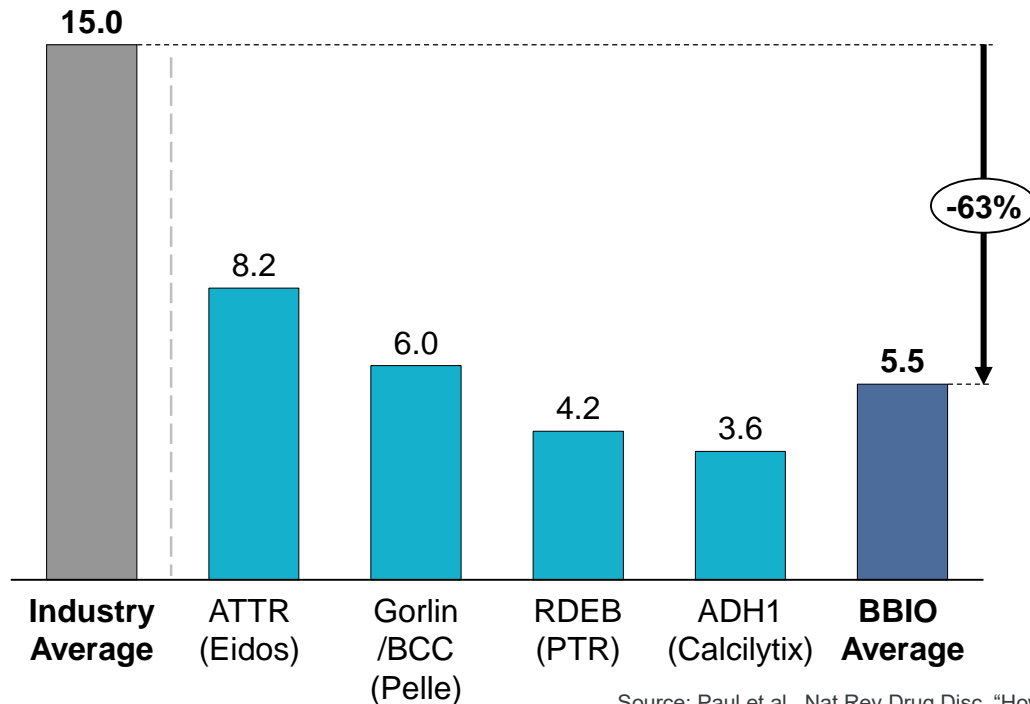
Spend to IND

# On average, we have brought assets forward more efficiently than industry average

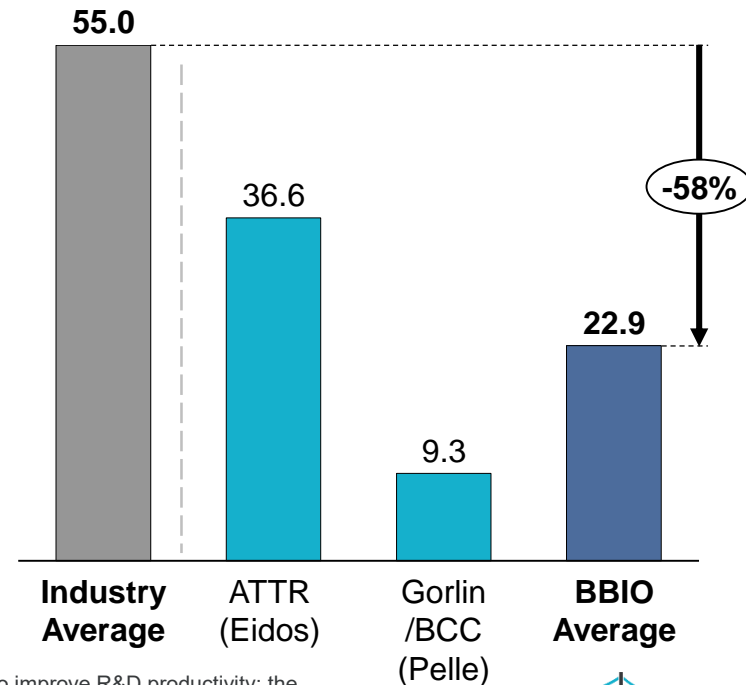
## Operationally Efficient Platform

- Our track record to date is ~\$6mm to IND and is ~\$23mm IND to POC (Phase 2)
- We aim to **rapidly and decisively** advance our product candidates to objective critical decision points
- We field a minimum viable team for each asset, with the goal of ensuring that each program has sufficient personnel to fit its purpose while **reducing excess overhead cost**

## Spend to IND (\$mm)



## Spend IND to POC (\$mm)



Note: BBIO values exclude license and acquisition costs.

Source: Paul et al., Nat Rev Drug Disc, "How to improve R&D productivity: the pharmaceutical industry's grand challenge.", 2010

# 2019 included a range of accomplishments across our development programs and operations

## Clinical, regulatory, and scientific

- ✓ **ATTR:** initiated Ph3 trial, presented Ph2 open-label extension data
- ✓ **Achondroplasia:** initiated Ph2 observational lead-in, established therapeutic window between human safety database and projected efficacious achondroplasia doses
- ✓ **RDEB:** initiated Ph2 POC trial
- ✓ **MOCD Type A:** initiated rolling NDA
- ✓ **BCC:** completed Gorlin Ph3 enrollment; initiated Ph2 high frequency BCC
- ✓ **Oncology:** SHP2 combination data w/ MEK, EGFR, KRAS augmenting inhibition; GPX4 demonstrated monotherapy activity in mouse model
- ✓ **CCA:** fast track designation for first line, completed enrollment in second line efficacy cohort
- ✓ **CAH:** demonstrated 6-month durability in adrenal cortex
- ✓ **Canavan:** demonstrated broad CNS distribution using IV route of administration

## Operations and finance

- ✓ **Building commercial organization:** Jennifer Cook, BOD member and commercial advisor; appointed Matt Outten as CCO
- ✓ **Financing:** raised over \$650M in IPO and private financing

# BridgeBio: Commercial build out

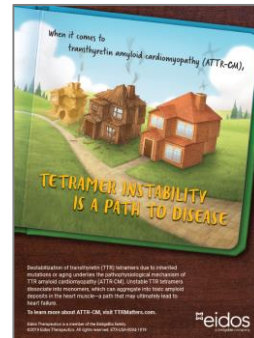
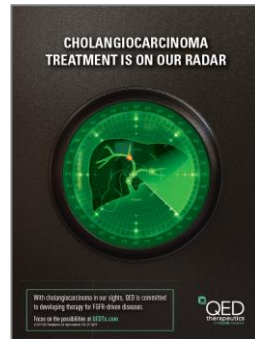
## Top talent makes a difference

### • CCO: Matt Outten



- 20+ years pharma/biotech
- Multiple commercial leadership positions across sales, marketing, market access
- Led the successful launch of Imbruvica for 6 indications
- **25 BBIO leadership roles:**
  - VPs of Marketing, Market Access, Distribution, Commercial Operations, Directors of Marketing and Training, Data Analytics and Operations
  - In-field teams established: Clinical Trial Liaisons, Professional Services

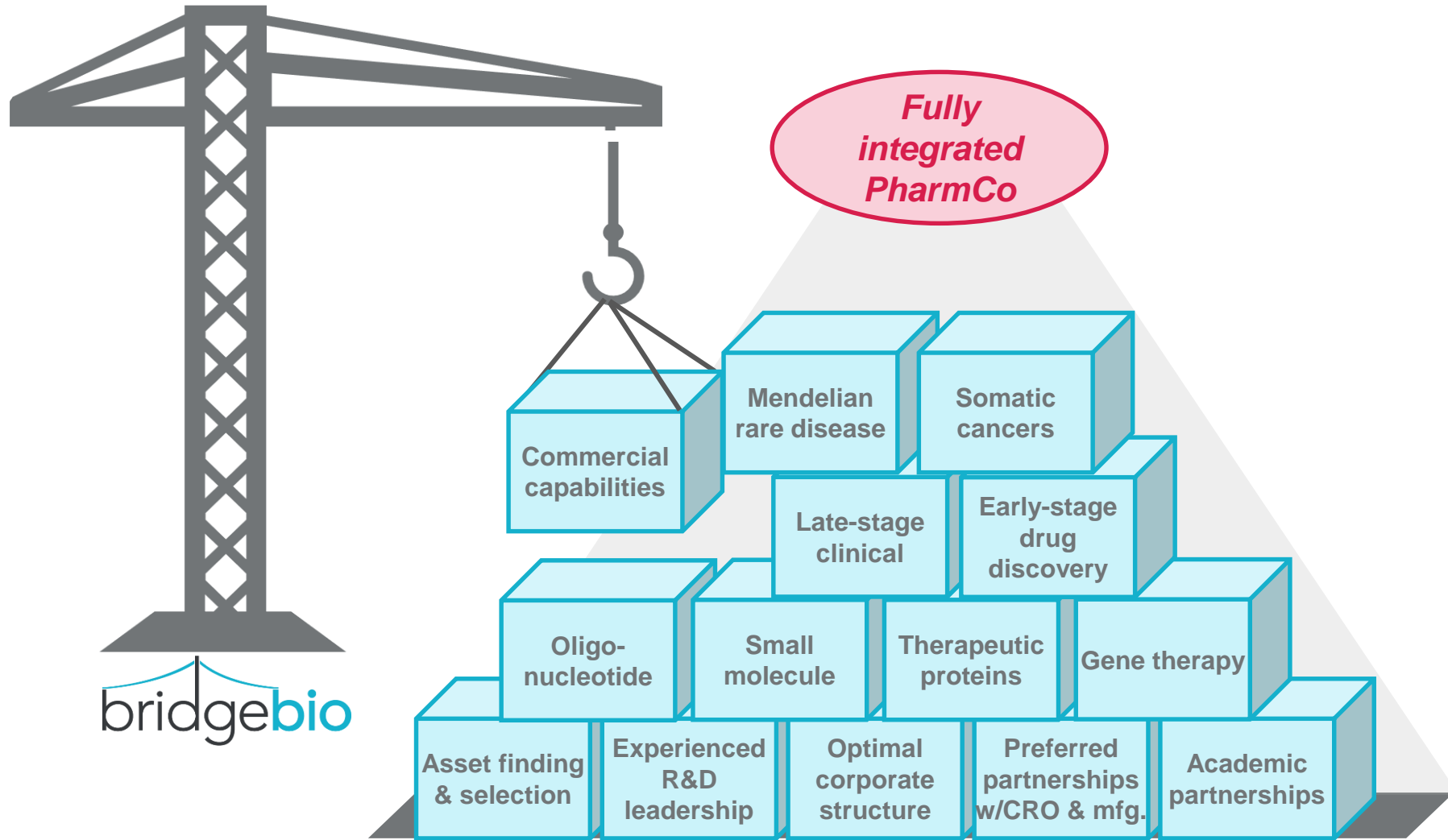
## Building awareness



## Gearing up for genetic disease launches

- Critical **patient identification capabilities expanded** in rare diseases with multiple data sources
- Allows better target planning, asset review, and appropriate resource allocation
- Developing **best in class HUB and patient assistance programs** in prep for commercial launches
- **Developing tailored launch plans** for each sub, from brand development to promotional material and in-field team training

# We are building a leading integrated pharma company



# Multiple catalysts anticipated in 2020-2021

ESTIMATED

2020		2021
1H	2H	FY
<ul style="list-style-type: none"> <li>✓ New program announcements</li> <li>✓ FGFRi for cancer: FPI Ph3 adjuvant urothelial carcinoma study</li> <li>✓ FGFRi for cancer: FPI Ph2 FGFR fusion tumor agnostic Ph2 study</li> </ul>	<ul style="list-style-type: none"> <li>❑ <b>Recombinant COL7 for RDEB: Topline Ph1/2 data</b></li> <li>❑ FGFRi for cancer: Pivotal 2L CCA data</li> <li>❑ <b>Low-dose FGFRi for achondroplasia: Begin dosing Ph2</b></li> <li>❑ <b>TTR stabilizer for ATTR: Complete enrollment of ATTR-CM Ph3</b></li> <li>❑ FGFRi for cancer: Submit NDA for 2L CCA</li> <li>❑ cPMP for MoCD type A: Complete NDA submission</li> <li>❑ Multiple new IND filings</li> </ul>	<ul style="list-style-type: none"> <li>❑ <b>TTR stabilizer for ATTR: Topline data Ph3 Part A in ATTR-CM</b></li> <li>❑ <b>Topical SMOi for Gorlin: Topline Ph3 data</b></li> <li>❑ <b>Low-dose FGFRi for achondroplasia: Ph2 PoC data</b></li> <li>❑ <b>CAH gene therapy: Ph1/2 PoC data</b></li> <li>❑ <b>FGFRi for cancer: 2L CCA approval and launch</b></li> <li>❑ <b>cPMP for MoCD type A: Approval and launch</b></li> <li>❑ <b>CaSR antagonist for ADH1: Ph2 POC data</b></li> </ul>

**Strong balance sheet expected to provide runway into 2022:**

**\$577mn as of YE19 plus \$550mn in gross proceeds from recent convertible debt offering**