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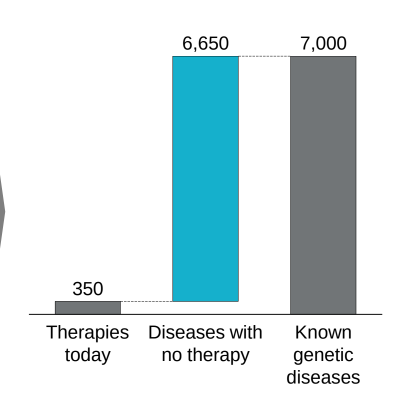


We are at Day 1 in the era of genetic medicine

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	PORTOLA' Globalblood THERAPEUTICS MyoKardia MILLENNIUM THE MAETA OKCOLOGY COMPANY	VELCADE* (bortezomib) ron nucernov Coopleterindov /o gecombert/ inchided-zho Coopleterindov /o gecombert/ inchided-zho Coopleterindov /o coopleterindov /			
Frank McCormick, PhD, FRS Chairman of Oncology	ONYX UCSF	(sorafenib) tablets Kyprolis (carfilzomib) Specton			
Richard Scheller, PhD Chairman of R&D	Genentech 23andMe	TECENTRIQ alezolizumab PERJETA pertuzumab insulati OCREVUS OC			

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



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

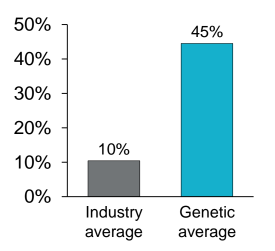


We believe genetic disease drug discovery is lower risk, faster, with potentially higher returns than traditional drug discovery

>4x

Higher cumulative probability of success

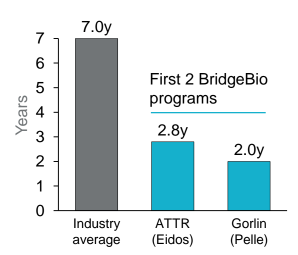
Probability of success from Ph1 to launch



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



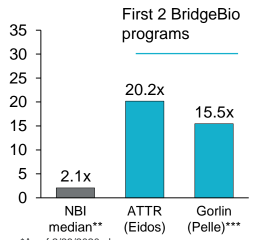
Sources: Paul et al., Nat Rev Drug Disc, "How to improve R&D productivity: the pharmaceutical industry's grand challenge.", 2010



Better return on investment*

Total return on investment

[Enerprise 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

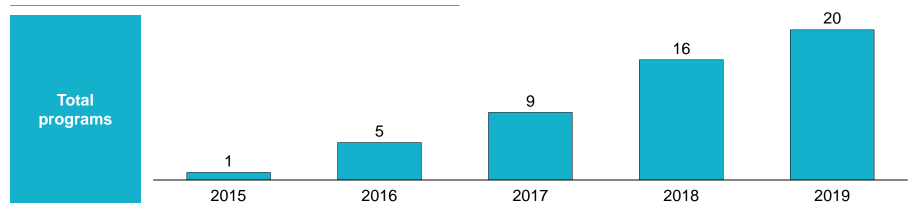




A rapidly-advancing pipeline

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:



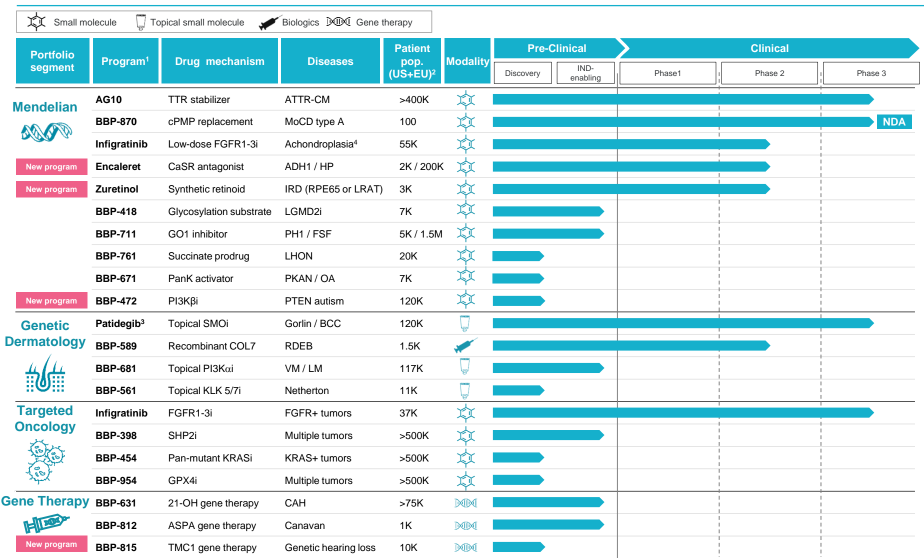
Advancement of product candidates through key stage-gates:

Internal program advancement

	Lead-Opt	IND enabling	Ph1	Ph2	Ph3
2016		+1			
2017			→ +1		
2018		+1		→ +1	
2019		+6	→ +1		+3



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



¹ Each of our programs is housed in a separate subsidiary; ² Patient population: Prevalence except for asterisked figures which represent incidence; ³We 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. ⁴Protocol accepted by Australian local ethics committed, 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

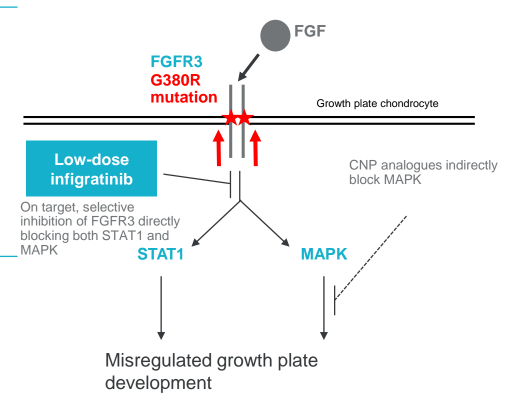
Potential best-in-class approach to treating achondroplasia directly at its genetic source

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

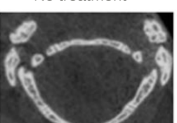
1 Cranial bone issues

17% increase in FM area

6% increase in AP skull length

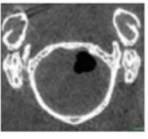
May lead to **decrease** in **foramen magnum stenosis** and fewer surgeries

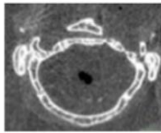
FGFR3 WT No treatment



FGFR3^{Y367C/+}
No treatment

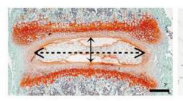




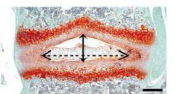


2 Disorders of the spine

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

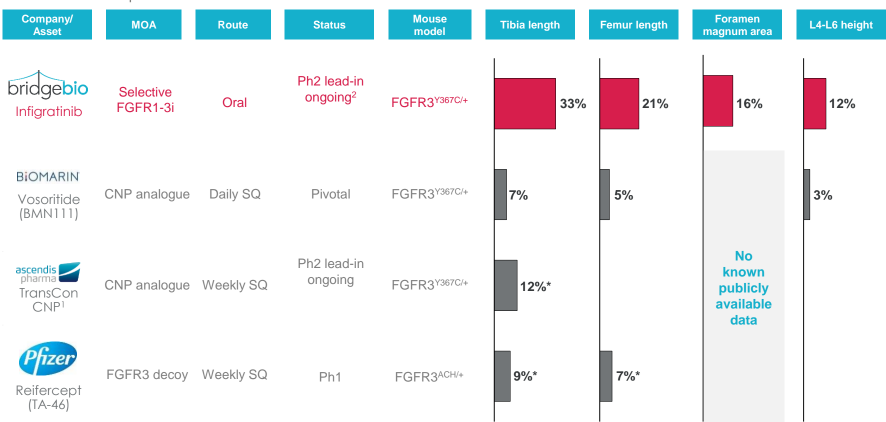
Note: percent increase compared to vehicle treated FGFR3Y367C/+ mouse, infigratinib treatment with 2mg/kg subcutaneous dose



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



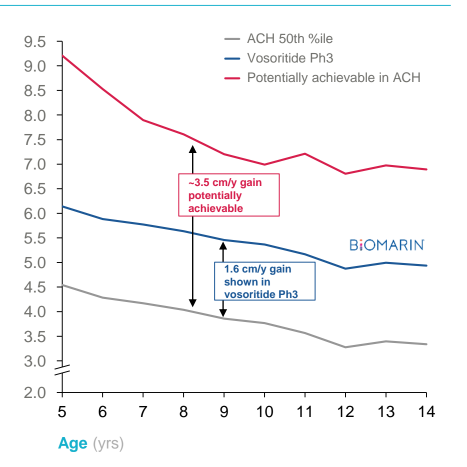
Source: Komla-Ebri et al. J Clin In2v 2016, Lorget et al. Am J Hum Genet 2012, Garcia et al. Science Trans Med 2013, Breinholt ENDO 2017 Note: subcutaneous doses, percent increase compared to vehicle treated FGFR3^{Y367C/+}, FGFR3^{ACH/+} mouse as noted in "Mouse model" columns Infigratinib treatment with 2mg/kg subcutaneous dose

¹Based on vosoritide continuous infusion; *Value estimated using Digitizelt. ²Protocol submitted to Australian local ethics committed, IND submission to FDA expected 2020.



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



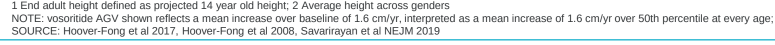


End adult height in feet^{1,2} Projected height using AGV

5 4 77 Potential end adult height with treatment that increases AGV to 95th %ile of average stature from ages 5-14

Projected end adult height for children treated with vosoritide from ages 5-14 without tachyphylaxis

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



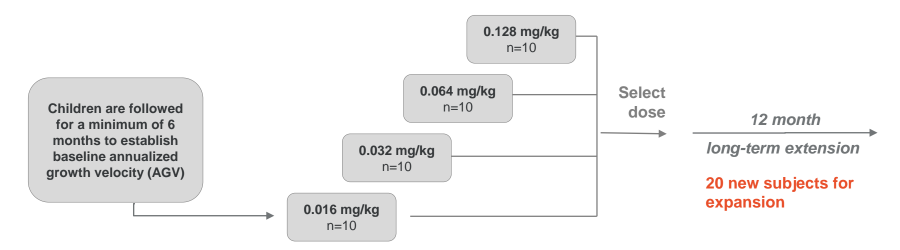


The PROPEL clinical program is enrolling and potential POC data expected in 2021

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)



Bardy, child with 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 dermalepithelial 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

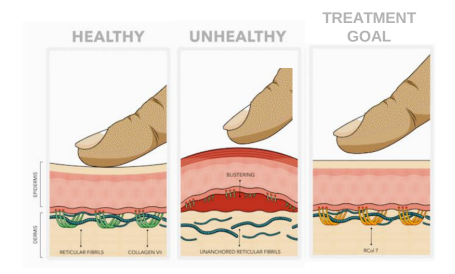
Recombinant collagen type VII for recessive dystrophic epidermolysis bullosa (RDEB)

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

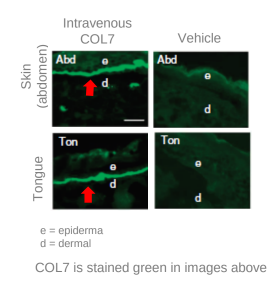


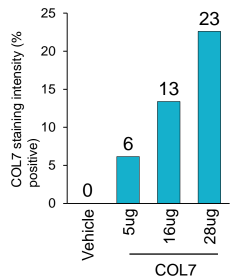


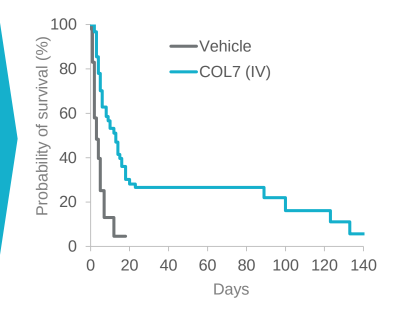
Recombinant COL7 distributes systemically, leading to survival benefits in the RDEB mouse model

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





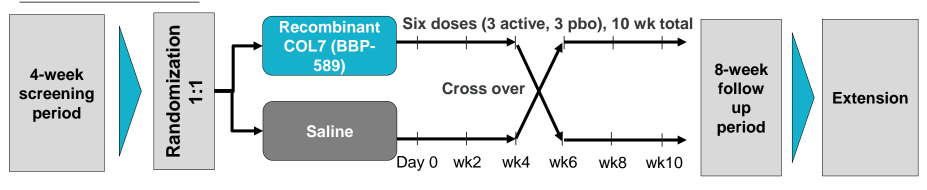




Ongoing randomized, dose-escalation Phase 1/2 proof-of-concept clinical study in adults with RDEB

- First patient dosed in 1Q19
- Anticipate potential POC data in 2020

Protocol for EACH cohort



Cohort 1: 0.1 mg/Kg N=2 Cohort 2: 0.3 mg/Kg N=4 Cohort 3: 1 mg/Kg N=6

Cohort 4: 3 mg/Kg N=4

KEY INCLUSION CRITERIA

- Adult with RDEB diagnosis
- Deficiency but not total loss of COL7 protein
- At least 1 wound >20cm² 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



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

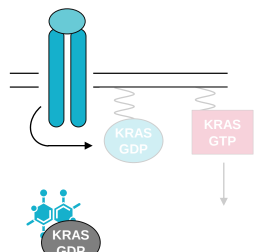
KRAS activation in cancer is a multistep process

Receptor tyrosine kinase signals (EGFR, FGFR, etc) **KRAS GTP** KRAS is activated by RTK signaling KRAS KRAS must tether to Active KRAS the cell membrane to drives cancer be activated proliferation and survival

Our programs target different steps of the KRAS activation process

Program 1: C185 targeting

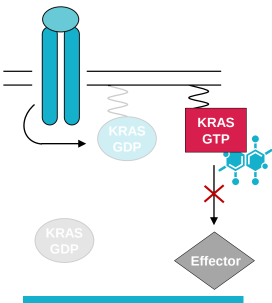
- · Blocks KRAS from tethering
- Blocks conversion of inactive KRAS GDP to active KRAS GTP



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

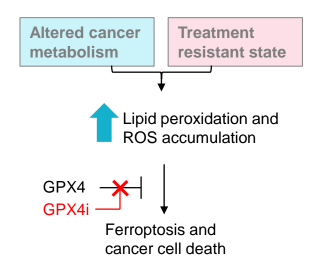
In vitro properties BBIO RVMD* pERK IC ₅₀ (nM) cellular assay <40 <40 hERG Patch clamp IC ₅₀ (μM) >100 ? Monotherapy anti-tumor activity KRASG12C xenograft √ √ EGFR mutant xenograft √ √ Combination enhanced anti-tumor activity G12Ci √(AMG 510) √(MRTX 849) MEKi √(trametinib) √(cobimetinib) EGFRi osimertinib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

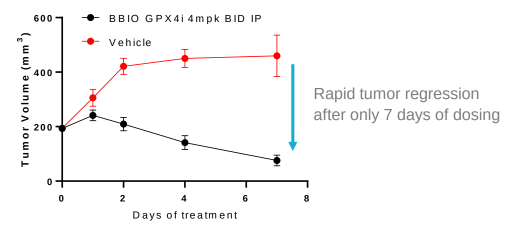


- GPX4 was recently identified as a key tumor dependency in multiple CRISPR screens and mechanistic studies
- GPX4 allows tumor cells to survive by neutralizing toxic lipid peroxides
- Our approach is to directly inhibit GPX4, thereby triggering cancer death through ferroptosis

Key data

In vivo monotherapy activity in a renal cell carcinoma mouse model

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



Synergy with targeted therapies and immunotherapy using in vitro models



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

FGFR2+
cholangiocarcinoma
3 1 2 3 3

Indication

Key Data

Status

Next planned update

- **39% ORR**
- in patients with ≤1 previous line of treatment
- Enrollment complete in 2L Ph2 pivotal cohort
- Ph3 in 1L study enrolling
- Updated pivotal data 2H20
- NDA submission 2H20
- 2021 launch

FGFR3+ urothelial carcinoma

25% ORR

in metastatic relapsed refractory setting suggests clear activity in this tumor

- FPI for Ph3 in adjuvant setting in 1H20
- Complete enrollment in Ph3 adjuvant study

FGFR fusionpositive tumor agnostic

5 tumor types

Showed response to infigratinib in Ph1/2

- FPI for Ph2 signal optimization study in 1H20
- Potential Ph2 data 2021



Gene therapy portfolio



Experienced team with track record in gene therapy









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

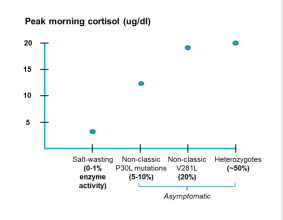
TMC1-driven hearing loss (BBP-815)

- 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

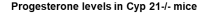
21-OH gene therapy for CAH: NHP study showed durable transgene expression; 5-10% of WT enzyme may be sufficient for clinical impact

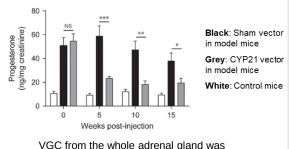
Genotype-phenotype studies show that >5-10% of enzyme activity results in nonclassical CAH



 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

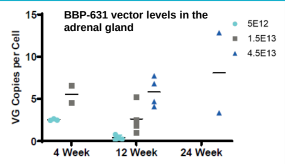




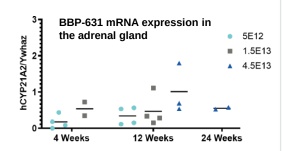
 At 15 weeks in treated mice, progesterone (the key substrate of 210Hase in mice) was significantly reduced vs untreated mice

0.13 at wk 18

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 dosedependent and stable out at 24 wks

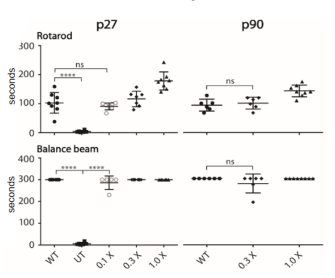




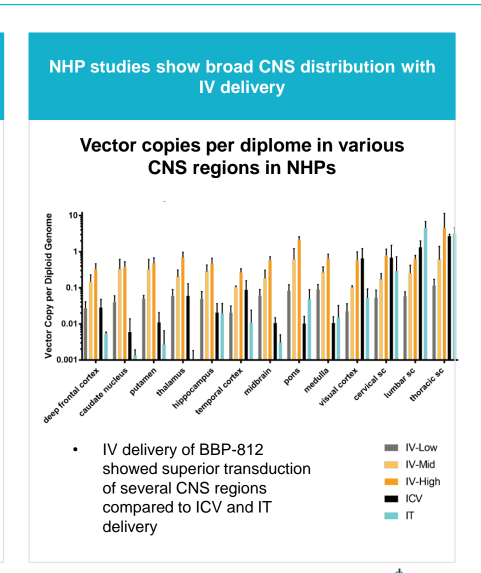
ASPA gene therapy for Canavan: Phenotypic correction in a lethal mouse model and broad CNS transduction in NHPs

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



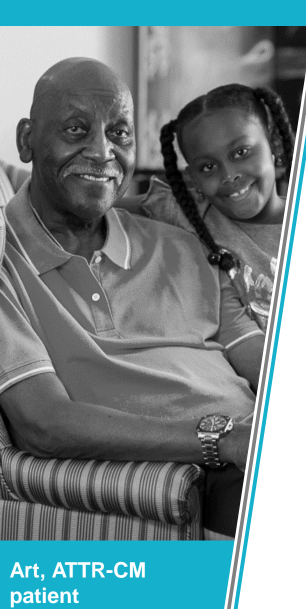
ASPA KO mice treated with at least 2.6e13
vg/kg had NAA metabolism and performance
on motor function tests restored. Mice treated
at 2.6e14 vg/kg outperformed WT mice.



Source: ESGCT 2019

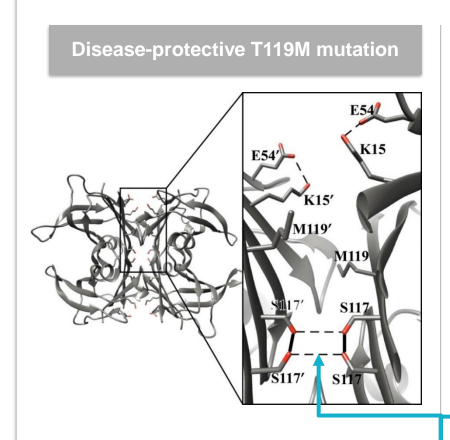


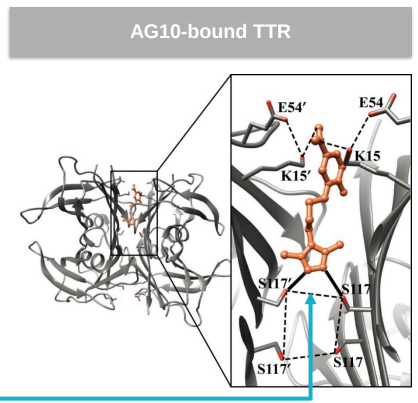
AG10 for TTR amyloidosis (Eidos)



- 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 topline data expected in 2021

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

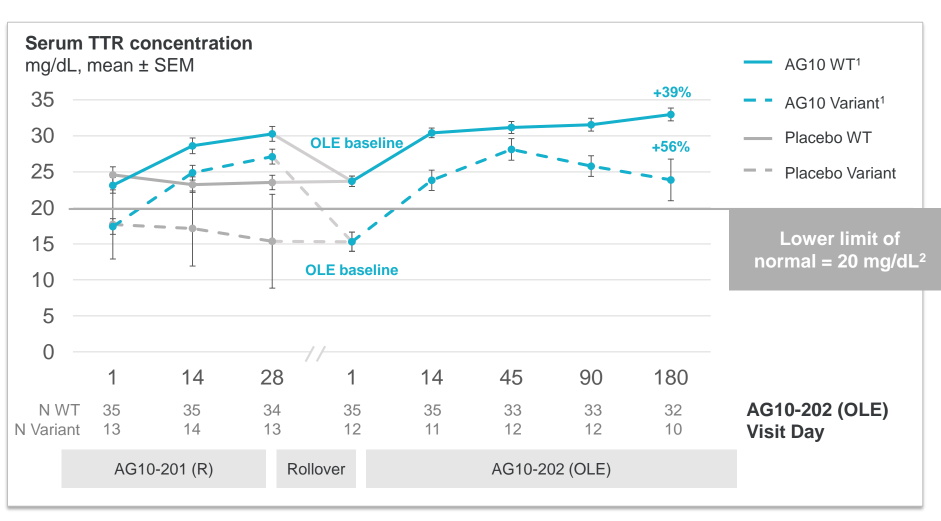




Strong inter-monomer H-bonds observed via X-ray crystallography Unique binding mode vs other stabilizers



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

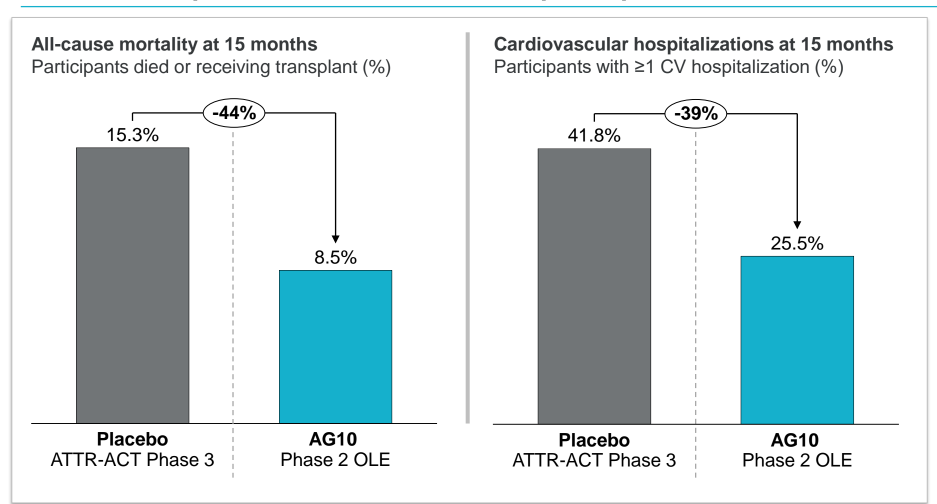


^{1 400}mg and 800mg BID AG10 groups pooled during randomized portion



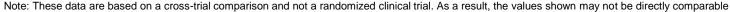
² Defined as the lower limit of the reference interval for the serum prealbumin (TTR) clinical laboratory assay

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



Phase 3 ATTRibute study expected to complete enrollment in 2H20

¹ Based on routine adverse event reporting





Fosdenopterin (cPMP replacement) for MoCD type A



Genetic driver: MOCS1 / cPMP depletion

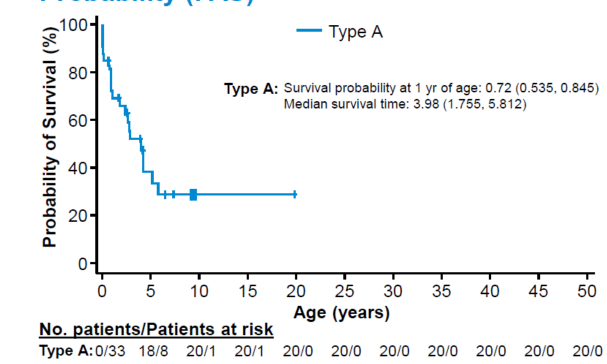
Prevalence (US + EU): 100

- 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

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)

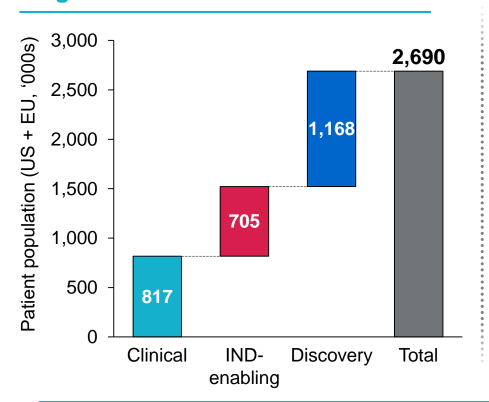


Note: Kaplan-Meier curves step down at time points at which a death has been observed; slashes represent patients whose observation time was censored as of the last contact date.



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

Total:	817.000
MoCD type A	100
RDEB	1,500
Inherited retinal dystrophy	3,000
FGFR+ cancer	37,000
Achondroplasia	55,000
Basal cell carcinoma	120,000
Hypoparathyroidism	200,000
Indication ATTR	Population 400,000

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

Zuretinol

BBP-815



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: PI3Kβ inhibitor
Diseases and prevalence:

PTEN autism
120,000
US + EU

Mechanism: TMC1 gene therapy Diseases and prevalence:

Genetic hearing loss 10,000 US + EU

Modality: Small molecule

Discovery

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

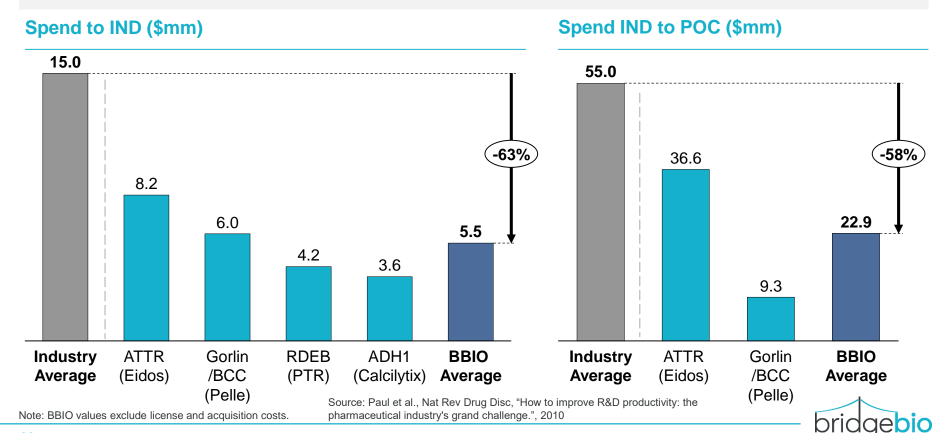
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On average, we have brought assets forward more efficiently than industry average

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



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





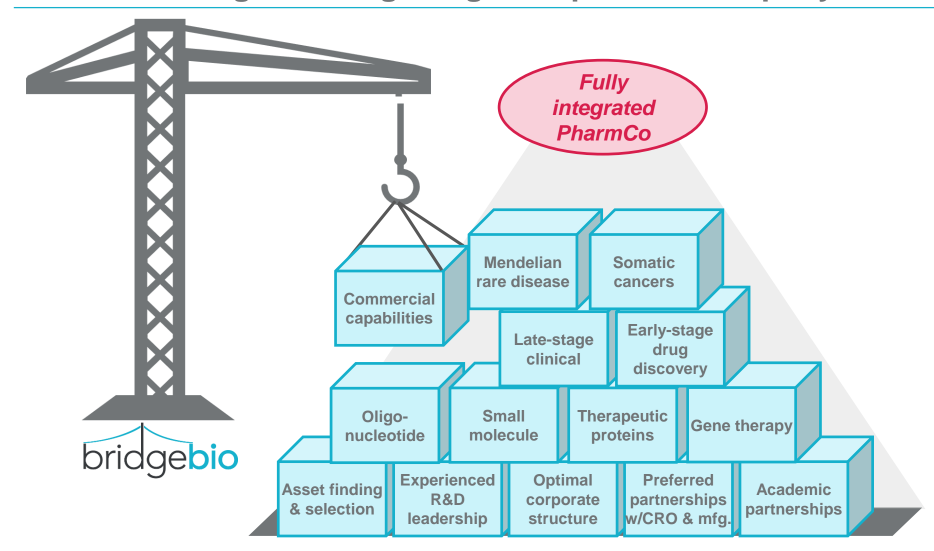


Gearing up for genetic disease launches

- 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



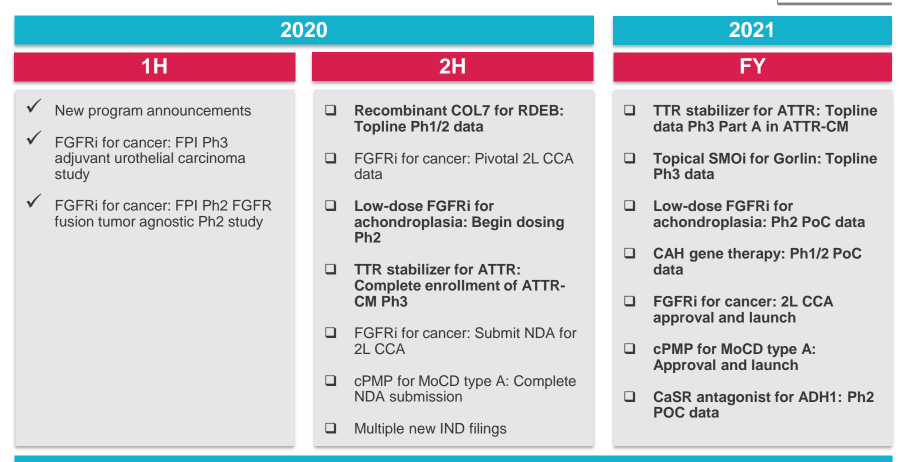
We are building a leading integrated pharma company





Multiple catalysts anticipated in 2020-2021

ESTIMATED



Strong balance sheet expected to provide runway into 2022: \$577mn as of YE19 plus \$550mn in gross proceeds from recent convertible debt offering

