

UNITED STATES
SECURITIES AND EXCHANGE COMMISSION
WASHINGTON, D.C. 20549

FORM 8-K

CURRENT REPORT

Pursuant to Section 13 or 15(d) of the Securities Exchange Act of 1934

Date of Report (Date of earliest event reported): January 13, 2025

BridgeBio Pharma, Inc.

(Exact name of Registrant as Specified in Its Charter)

Delaware
(State or Other Jurisdiction of Incorporation)

001-38959
(Commission File Number)

84-1850815
(IRS Employer Identification No.)

3160 Porter Dr., Suite 250
Palo Alto, CA
(Address of Principal Executive Offices)

94304
(Zip Code)

Registrant's Telephone Number, Including Area Code: (650) 391-9740

Not Applicable
(Former Name or Former Address, if Changed Since Last Report)

Check the appropriate box below if the Form 8-K filing is intended to simultaneously satisfy the filing obligation of the registrant under any of the following provisions:

- Written communications pursuant to Rule 425 under the Securities Act (17 CFR 230.425)
- Soliciting material pursuant to Rule 14a-12 under the Exchange Act (17 CFR 240.14a-12)
- Pre-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))
- Pre-commencement communications pursuant to Rule 13e-4(c) under the Exchange Act (17 CFR 240.13e-4(c))

Securities registered pursuant to Section 12(b) of the Act:

Title of each class	Trading Symbol(s)	Name of each exchange on which registered
Common Stock, par value \$0.001 per share	BBIO	The Nasdaq Global Select Market

Indicate by check mark whether the registrant is an emerging growth company as defined in Rule 405 of the Securities Act of 1933 (§ 230.405 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (§ 240.12b-2 of this chapter).

Emerging growth company

If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act.

Item 2.02 Results of Operations and Financial Condition.

On January 13, 2025, BridgeBio Pharma, Inc., or the Company, issued a press release that contains certain preliminary financial information as of and for the fiscal year ended December 31, 2024. Specifically, the press release states that the Company received \$500 million upon acoramidis U.S. Food and Drug Administration approval.

The information in this Item 2.02 is unaudited and preliminary, and does not present all information necessary for an understanding of the Company's results of operations for the fiscal year ended December 31, 2024, or financial condition as of December 31, 2024. The audit of the Company's financial statements for the year ended December 31, 2024 is ongoing and could result in changes to the information in this Item 2.02.

Item 7.01 Regulation FD Disclosure.

The disclosure in Item 2.02 above is hereby incorporated by reference into this Item 7.01, and a copy of the press release referenced in Item 2.02 is furnished as Exhibit 99.1 hereto.

The information contained in Items 2.02 and 7.01, as well as Exhibit 99.1, to this Current Report on Form 8-K shall not be deemed to be "filed" for purposes of Section 18 of the Securities Exchange Act of 1934, as amended (the "Exchange Act"), or otherwise subject to the liabilities of that section, nor shall such information be deemed incorporated by reference in any filing under the Securities Act of 1933, as amended, or the Exchange Act, except as expressly set forth by specific reference in such a filing.

Item 8.01 Other Events.

On January 13, 2025, the Company also presented a business update at the 43rd Annual J.P. Morgan Healthcare Conference. A copy of the Company's presentation slides, which has been published on the Company's website, is filed as Exhibit 99.2 to this current report on Form 8-K and is incorporated by reference herein.

Forward Looking Statements

This Current Report on Form 8-K contains forward-looking statements. Statements in this Current Report on Form 8-K or the materials furnished or filed herewith may include statements that are not historical facts and are considered forward-looking within the meaning of Section 27A of the Securities Act of 1933, as amended (the Securities Act), and Section 21E of the Securities Exchange Act of 1934, as amended (the Exchange Act), which are usually identified by the use of words such as "anticipates," "believes," "continues," "estimates," "expects," "hopes," "intends," "may," "plans," "projects," "remains," "seeks," "should," "will," and variations of such words or similar expressions. The Company intends these forward-looking statements to be covered by the safe harbor provisions for forward-looking statements contained in Section 27A of the Securities Act and Section 21E of the Exchange Act. These forward-looking statements, including express and implied statements relating to the Company's expectations regarding the commercial success of Attruby; the Company's clinical trials, including the timing of the last patient-last visit and topline data readouts for each of FORTIFY, CALIBRATE, and PROPEL 3; the potential for encaletet to become a new treatment for ADHI; the potential for BBP-418 to become a new treatment for LGMD2I/R9; the potential for infgratinib to become a new treatment for achondroplasia; timing of approval of Attruby for ATTR-CM in the European Union and Japan; and the Company's preliminary and unaudited estimate of cash and the Company's anticipated funding of its current operations and related timelines; and the Company's expectations regarding reaching regulatory milestones and receipt of milestone payments, among others, reflect the Company's current views about the Company's plans, intentions, expectations and strategies, which are based on the information currently available to us and on assumptions the Company has made. Although the Company believes that its plans, intentions, expectations and strategies as reflected in or suggested by those forward-looking statements are reasonable, the Company can give no assurance that the plans, intentions, expectations or strategies will be attained or achieved. Furthermore, actual results may differ materially from those described in the forward-looking statements and will be affected by a number of risks, uncertainties and assumptions, including, but not limited to, initial and ongoing data from the Company's preclinical studies and clinical trials not being indicative of final data, the potential size of the target patient populations the Company's product candidates are designed to treat not being as large as anticipated, the design and success of ongoing and planned clinical trials, future regulatory filings, approvals and/or sales, despite having ongoing and future interactions with the FDA or other regulatory agencies to discuss potential paths to registration for the Company's product candidates, the FDA or such other regulatory agencies not agreeing with the Company's regulatory approval strategies, components of the Company's filings, such as clinical trial designs, conduct and methodologies, or the sufficiency of data submitted, the continuing success of the Company's collaborations, the Company's ability to obtain additional funding, including through less dilutive sources of capital than equity financings, potential volatility in the Company's share price, the impacts of current macroeconomic and geopolitical events, including changing conditions from hostilities in Ukraine and in Israel and the Gaza Strip, increasing rates of inflation and changing interest rates, on business operations and expectations, as well as those risks set forth in the Risk Factors section of the Company's most recent Quarterly Report on Form 10-Q and Annual Report on Form 10-K and the Company's other filings with the U.S. Securities and Exchange Commission. Moreover, the Company operates in a very competitive and rapidly changing environment in which new risks emerge from time to time. These forward-looking statements are based upon the current expectations and beliefs of the Company's management as of the date of this Current Report on Form 8-K, and are subject to certain risks and uncertainties that could cause actual results to differ materially from those described in the forward-looking statements. Except as required by applicable law, the Company assumes no obligation to update publicly any forward-looking statements, whether as a result of new information, future events or otherwise.

(d) Exhibits.

Exhibit	Description
99.1	Press Release dated January 13, 2025.
99.2	Slides from BridgeBio Pharma, Inc.'s J.P. Morgan Healthcare Conference Presentation, dated January 13, 2025.
104	Cover Page Interactive Data File (embedded within the Inline XBRL document)

SIGNATURES

Pursuant to the requirements of the Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned hereunto duly authorized.

BridgeBio Pharma, Inc.

Date: January 13, 2025

By: /s/ Brian C. Stephenson
Brian C. Stephenson
Chief Financial Officer

BridgeBio Announces Commercial Progress, Program Updates, and 2025 Milestones

- Remarkable early Attruby demand: 430 scripts written by 248 unique HCPs since FDA approval with broad uptake across academic centers and community centers in all patient types

- Fully enrolled three major market Phase 3 clinical trials: FORTIFY (BBP-418 for LGMD2I/R9); CALIBRATE (encaleret for ADHI); and PROPEL 3 (infigratinib for Achondroplasia)

- Well-financed to launch Attruby and read out major market Phase 3 trials: \$406M in cash as of last quarter, received \$500M upon acoramidis FDA approval from royalty facility, and anticipate \$105M in regulatory milestones in 1H 2025 from acoramidis Europe and Japan approvals

PALO ALTO, CA – January 13, 2025 – BridgeBio Pharma, Inc. (Nasdaq: BBIO) (“BridgeBio” or the “Company”), a new type of biopharmaceutical company focused on genetic diseases, today provided updates on its commercial progress for Attruby (acoramidis), status of late-stage pipeline programs, and anticipated 2025 milestones.

“With the FDA’s approval of Attruby, we marked an important moment for both our organization and the broader ATTR-CM patient community in need of new treatment options. We’re grateful for the enthusiasm surrounding the product and the associated initial commercial momentum, with 430 prescriptions written by 248 unique physicians, and we look forward to continued progress,” said Neil Kumar, Ph.D., Founder and CEO of BridgeBio. “Additionally, we are excited to share that we have completed enrollment of all three of our major market Phase 3 clinical trials. I look forward to continuing to work with this stellar team to serve patients with genetic disease in 2025.”

Business Update

On November 22, 2024, the U.S. Food and Drug Administration (FDA) approved Attruby (acoramidis), a near-complete TTR stabilizer (≥90%), to reduce cardiovascular death and cardiovascular-related hospitalization in adult patients with ATTR-CM, a progressive fatal disease presenting as an infiltrative, restrictive cardiomyopathy resulting in heart failure.

Since the approval, BridgeBio has seen remarkable momentum with 430 patient prescriptions written by 248 physicians.

Pipeline Updates

BBP-418 – Glycosylation substrate for limb-girdle muscular dystrophy type 2I/R9 (LGMD2I/R9):

- FORTIFY is a Phase 3 clinical trial of BBP-418 in LGMD2I/R9, a rare genetic disorder caused by variants in the fukutin-related protein (FKRP) gene that result in progressive muscle degeneration and damage, and eventual loss of functional independence. The trial is fully enrolled with 112 patients.
-

- The Company expects Last Patient – Last Visit (LPLV) and topline readout of the interim analysis cohort in second half 2025.
- If successful, BBP-418 would be the first approved therapy for individuals living with LGMD2I/R9.

Encalereet – Calcium-sensing receptor (CaSR) antagonist for autosomal dominant hypocalcemia type 1 (ADH1):

- CALIBRATE, the Phase 3 clinical trial of encalereet in ADH1, a rare, genetic form of hypoparathyroidism, is fully enrolled with 70 patients. The trial is designed to evaluate the efficacy and safety of encalereet compared to standard of care in adult patients with ADH1.
- The Company expects Last Patient – Last Visit and topline readout in second half 2025.
- If successful, encalereet would be the first approved therapy for individuals living with ADH1.

Infigratinib – FGFR1-3 inhibitor for achondroplasia and hypochondroplasia:

- PROPEL 3, the Phase 3 clinical trial of infigratinib in achondroplasia, the most common form of disproportionate short stature, is fully enrolled with 114 participants.
- The Company expects Last Participant – Last Visit in second half 2025.
- If successful, infigratinib would be the first approved oral therapy for children living with achondroplasia.

2025 Milestones

Program	Status	Anticipated 2025 Milestone
Acoramidis for ATTR-CM	US FDA approval on November 22, 2024	EU and Japan approvals in 1H 2024
BBP-418 for LGMD2I/R9	FORTIFY, Phase 3 study enrollment completed	Last Patient – Last Visit and Topline readout in 2H 2025
Encalereet for ADH1	CALIBRATE, Phase 3 study enrollment completed	Last Patient – Last Visit and Topline readout in 2H 2025
Infigratinib for achondroplasia	PROPEL 3, Phase 3 study enrollment completed	Last Participant – Last Visit in 2H 2025

About Attruby™ (acoramidis)

INDICATION

Attruby is a transthyretin stabilizer indicated for the treatment of the cardiomyopathy of wild-type or variant transthyretin-mediated amyloidosis (ATTR-CM) in adults to reduce cardiovascular death and cardiovascular-related hospitalization.

IMPORTANT SAFETY INFORMATION

Adverse Reactions

Diarrhea (11.6% vs 7.6%) and upper abdominal pain (5.5% vs 1.4%) were reported in patients treated with Attruby versus placebo, respectively. The majority of these adverse reactions were mild and resolved without drug discontinuation. Discontinuation rates due to adverse events were similar between patients treated with Attruby versus placebo (9.3% and 8.5%, respectively).

About BridgeBio Pharma, Inc.

BridgeBio Pharma, Inc. (BridgeBio) is a new type of biopharmaceutical company founded to discover, create, test, and deliver transformative medicines to treat patients who suffer from genetic diseases. BridgeBio's pipeline of development programs ranges from early science to advanced clinical trials. BridgeBio was founded in 2015 and its team of experienced drug discoverers, developers and innovators are committed to applying advances in genetic medicine to help patients as quickly as possible. For more information visit [bridgebio.com](https://www.bridgebio.com) and follow us on LinkedIn, Twitter and Facebook.

BridgeBio Pharma, Inc. Forward-Looking Statements

This press release contains forward-looking statements. Statements in this press release may include statements that are not historical facts and are considered forward-looking within the meaning of Section 27A of the Securities Act of 1933, as amended (the Securities Act), and Section 21E of the Securities Exchange Act of 1934, as amended (the Exchange Act), which are usually identified by the use of words such as “anticipates,” “believes,” “continues,” “estimates,” “expects,” “hopes,” “intends,” “may,” “plans,” “projects,” “remains,” “seeks,” “should,” “will,” and variations of such words or similar expressions. BridgeBio intends these forward-looking statements to be covered by the safe harbor provisions for forward-looking statements contained in Section 27A of the Securities Act and Section 21E of the Exchange Act. These forward-looking statements, including express and implied statements relating to the Company’s expectations regarding the commercial success of Attruby; the Company’s clinical trials, including the timing of the last patient-last visit and topline data readouts for each of FORTIFY, CALIBRATE and PROPEL 3; the potential for encaeteret to become a new treatment for ADHD1; the potential for BBP-418 to become a new treatment for LGMD2I/R9; the potential for infigratinib to become a new treatment for achondroplasia; timing of approval of Attruby for ATTR-CM in the European Union and Japan; and the Company’s preliminary and unaudited estimate of cash and the Company’s anticipated funding of its current operations and related timelines; and the Company’s expectations regarding reaching regulatory milestones and receipt of milestone payments, among others, reflect the Company’s current views about the Company’s plans, intentions, expectations and strategies, which are based on the information currently available to us and on assumptions the Company has made. Although the Company believes that its plans, intentions, expectations and strategies as reflected in or suggested by those forward-looking statements are reasonable, the Company can give no assurance that the plans, intentions, expectations or strategies will be attained or achieved. Furthermore, actual results may differ materially from those described in the forward-looking statements and will be affected by a number of risks, uncertainties and assumptions, including, but not limited to, initial and ongoing data from the Company’s preclinical studies and clinical trials not being indicative of final data, the potential size of the target patient populations the Company’s product candidates are designed to treat not being as large as anticipated, the design and success of ongoing and planned clinical trials, future regulatory filings, approvals and/or sales, despite having ongoing and future interactions with the FDA or other regulatory agencies to discuss potential paths to registration for the Company’s product candidates, the FDA or such other regulatory agencies not agreeing with the Company’s regulatory approval strategies, components of the Company’s filings, such as clinical trial designs, conduct and methodologies, or the sufficiency of data submitted, the continuing success of the Company’s collaborations, the Company’s ability to obtain additional funding, including through less dilutive sources of capital than equity financings, potential volatility in the Company’s share price, the impacts of current macroeconomic and geopolitical events, including changing conditions from hostilities in Ukraine and in Israel and the Gaza Strip, increasing rates of inflation and changing interest rates, on business operations and expectations, as well as those risks set forth in the Risk Factors section of the Company’s most recent Quarterly Report on Form 10-Q and Annual Report on Form 10-K and the Company’s other filings with the U.S. Securities and Exchange Commission. Moreover, the Company operates in a very competitive and rapidly changing environment in which new risks emerge from time to time. These forward-looking statements are based upon the current expectations and beliefs of the Company’s management as of the date of this press release, and are subject to certain risks and uncertainties that could cause actual results to differ materially from those described in the forward-looking statements. Except as required by applicable law, BridgeBio assumes no obligation to update publicly any forward-looking statements, whether as a result of new information, future events or otherwise.

BridgeBio Media Contact:

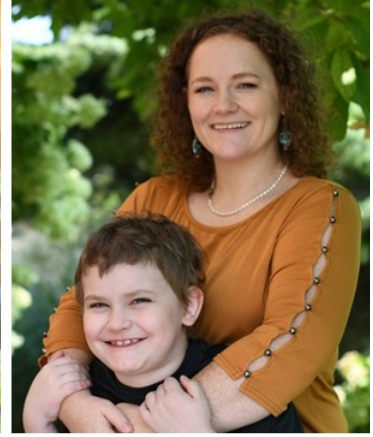
Bubba Murarka, EVP Communications
contact@bridgebio.com
(650)-789-8220

bridgebio

hope through
rigorous science

J.P. Morgan
Presentation

January 13, 2025




Forward Looking Statements and Disclaimer

The presentation contains forward-looking statements. Statements made or presented may include statements that are not historical facts and are considered forward-looking within the meaning of Section 27A of the Securities Act of 1933, as amended, and Section 21E of the Securities Exchange Act of 1934, as amended. Words such as “believe,” “anticipate,” “plan,” “expect,” “intend,” “will,” “may,” “goal,” “potential,” “should,” “could,” “aim,” “estimate,” “predict,” “continue” and similar expressions or the negative of these terms or other comparable terminology are intended to identify forward-looking statements, though not all forward-looking statements necessarily contain these identifying words. We intend these forward-looking statements to be covered by the safe harbor provisions for forward-looking statements contained in Section 27A of the Securities Act and Section 21E of the Exchange Act. These forward-looking statements, including express and implied statements relating to the commercial success of Attruby, the timing of ongoing clinical trials, including BridgeBio Oncology Therapeutics’ and Gondola Bio’s clinical trials, the clinical, therapeutic and market potential of our clinical development programs and our pipeline, BridgeBio Oncology Therapeutics’ pipeline and Gondola Bio’s pipeline, our speed of creating new and meaningful drugs and related impact on patients, the efficiency of our engine to rapidly and efficiently deliver medicines, our value creation potential for patients, the potential market sizes and opportunities, the safety, efficacy and mechanisms of our newly FDA-approved Attruby (acoramidis) and other later-stage products including infigratinib, BBP-418 and encaleret, the timing of approval of Attruby for ATTR-CM in the European Union and Japan, our financial position, including our expectations regarding reaching regulatory milestones and the receipt of milestone payments, the potency and safety of our product candidates, the potential benefits of our product candidates, the potential for greater patient access to medications, the affordability and availability of insurance coverage of our medications, and the timing and expectations regarding results of our various clinical trials, reflect our current views about our plans, intentions, expectations and strategies, which are based on the information currently available to us and on assumptions we have made. Such statements reflect the current views of the Company with respect to future events and are subject to known and unknown risks, including business, regulatory, economic and competitive risks, uncertainties, contingencies and assumptions about the Company, including, without limitation, risks inherent in developing therapeutic products, and those risks and uncertainties described under the heading “Risk Factors” in the Company’s most recent Annual Report on Form 10-K filed with the U.S. Securities and Exchange Commission (“SEC”) and in subsequent filings made by the Company with the SEC, which are available on the SEC’s website at www.sec.gov. In light of these risks and uncertainties, many of which are beyond the Company’s control, the events or circumstances referred to in the forward-looking statements, express or implied, may not occur. The actual results may vary from the anticipated results and the variations may be material. You are cautioned not to place undue reliance on these forward-looking statements, which speak to the Company’s current beliefs and expectations only as of the date of the presentation. Except as required by law, the Company disclaims any intention or responsibility for updating or revising any forward-looking statements made or presented at the presentation in the event of new information, future developments or otherwise. No representation is made as to the safety or effectiveness of the product candidates for the therapeutic use for which such product candidates are being studied.

Certain information communicated at the presentation may relate to or is based on studies, publications, surveys and other data obtained from third-party sources and the Company’s own internal estimates and research. While the Company believes these third-party sources to be reliable as of the date of the presentation, it has not independently verified, and makes no representation as to the adequacy, fairness, accuracy or completeness of, any information obtained from third-party sources. In addition, certain information to be communicated at the presentation involves a number of assumptions and limitations, and there can be no guarantee as to the accuracy or reliability of such assumptions. Finally, such research has not been verified by any independent source.

Such information is provided as of the date of the presentation and is subject to change without notice. The Company has not verified, and will not verify, any part of this presentation, and the Company makes no representation or warranty, express or implied, as to the accuracy or completeness of the information to be communicated at the presentation or as to the existence, substance or materiality of any information omitted from the presentation at the presentation. The Company disclaims any and all liability for any loss or damage (whether foreseeable or not) suffered or incurred by any person or entity as a result of anything contained or omitted from this document or the related presentation and such liability is expressly disclaimed.

Key recent achievements: clinical impact




The NEW ENGLAND
JOURNAL of MEDICINE

JANUARY 11, 2024

ORIGINAL ARTICLE

Efficacy and Safety of Acoramidis in Transthyretin Amyloid Cardiomyopathy

J.D. Gillmore, D.P. Judge, F. Cappelli, M. Fontana, P. Garcia-Pavia, S. Gibbs, M. Grogan, M. Hanna, J. Hoffman, A. Masri, M.S. Maurer, J. Nativi-Nicolau, L. Obici, S.H. Poulsen, F. Rockhold, K.B. Shah, P. Soman, J. Garg, K. Chiswell, H. Xu, X. Cao, T. Lystig, U. Sinha, and J.C. Fox, for the ATTRibute-CM Investigators*




The NEW ENGLAND
JOURNAL of MEDICINE

November 18, 2024

ORIGINAL ARTICLE

Oral Infigratinib Therapy in Children with Achondroplasia


R. Savarirayan, J.M. De Bergua, P. Arundel, J.P. Salles, V. Saraff, B. Delgado, A. Leva-Gea, H. McDevitt, M. Nicolino, M. Rossi, M. Salcedo, V. Cormier-Daire, M. Skae, P. Kannu, J. Phillips III, H. Saal, P. Harmatz, T. Candler, D. Hill, E. Muslimova, R. Weng, Y. Bai, S. Raj, J. Hoover-Fong, M. Irving, and D. Rogoff



The NEW ENGLAND
JOURNAL of MEDICINE

SEPTEMBER 28, 2023

CORRESPONDENCE



Efficacy and Safety of Encalceret in Autosomal Dominant Hypocalcemia Type 1

Rachel I. Gafni, M.D. Iris R. Hartley, M.D. Michael T. Collins, M.D.
National Institutes of Health



Trial enrolled **112 patients** to evaluate BBP-418 in **Limb-Girdle Muscular Dystrophy 21/R9**



Trial enrolled **70 participants** to evaluate encaleret in **Autosomal Dominant Hypocalcemia Type 1**



Trial enrolled **114 participants** to evaluate ifigratinib in **Achondroplasia**

Key recent achievements: regulatory advancement for late-stage programs



Infigratinib

Receives Breakthrough Designation (BTD) for Achondroplasia

BBP-418

Receives Rare Pediatric Disease Designation (RPDD) for Limb-Girdle Muscular Dystrophy 2I/R9

BBP-812

Receives Regenerative Medicine Advanced Therapy Designation (RMAT) for Canavan Disease

Key recent achievement: Attruby regulatory approval

 **Attruby**TM
(acoramidis) 356 mg tablet



Indicated for the treatment of adult patients with ATTR-CM to reduce cardiovascular death and cardiovascular-related hospitalization

430

Attruby Scripts

BridgeBio's growth is underpinned by an innovative corporate model, as highlighted in a recent publication



Applications of Portfolio Theory to Accelerating Biomedical Innovation

Neil Kumar, Andrew W. Lo, Chinmay Shukla, and Brian Stephenson

Diversified portfolio

- **Uncorrelated assets** enable multiple "shots on goal"
- **Modality and therapeutic area agnostic** selection of programs

Operational nimbleness

- **Centralized infrastructure** for functions with economies of scale
- **Hyper-focused and agile** decision making

World class, decentralized R&D

- Leverage **hyper-experienced**, science-focused R&D practitioners
- Asset-level incentives preserve **biology-level focus**

Creative financing toolkit

- Toolkit of **strategic options** (asset & portfolio levels)
- Costs are **variabilized and flexible** across central functions and affiliates

The BridgeBio ecosystem has a dynamic pipeline of products

Program	Indication	Pre-clinical	Phase 1	Phase 2	Phase 3	Approved	Patients (US + EU)	Market Oppty	Estimated US LOE**
Attruby (acoramidis)	Transthyretin Amyloidosis (ATTR-CM)					✓	500,000+	\$20B+	2039
Infigratinib	Achondroplasia (ACH)					Fully Enrolled	55,000+	\$2B+	2041
	Hypochondroplasia (HCH)						55,000+	\$2B+	2041
BBP-418	Limb-Girdle Muscular Dystrophy Type 2I/R9 (LGMD2I/R9)					Fully Enrolled	7,000+	\$1B+	2041
Encaleret	Autosomal Dominant Hypocalcemia Type 1 (ADH1)					Fully Enrolled	25,000+	\$1B+	2041
	Post-Surgical Hypoparathyroidism (PSH)						200,000+	\$1B+	2044
BBP-812	Canavan Disease				Phase 1/2 Pivotal		1,000	TBD	2039
BridgeBio Oncology Therapeutics	Oncology, various			38% ownership*			Various	Various	Various
GondolaBio	Rare disease, various			45% ownership*			Various	Various	Various

* BridgeBio Oncology Therapeutics and GondolaBio are separate, independent companies from BridgeBio. BridgeBio's initial interest in GondolaBio is subject to reduction as additional tranches of capital contributions are funded.
 ** Approximate estimated US LOE dates considering various factors including issued patents, pending patent applications, potential patent term extensions, and regulatory exclusivities.

NOW APPROVED



Attruby™

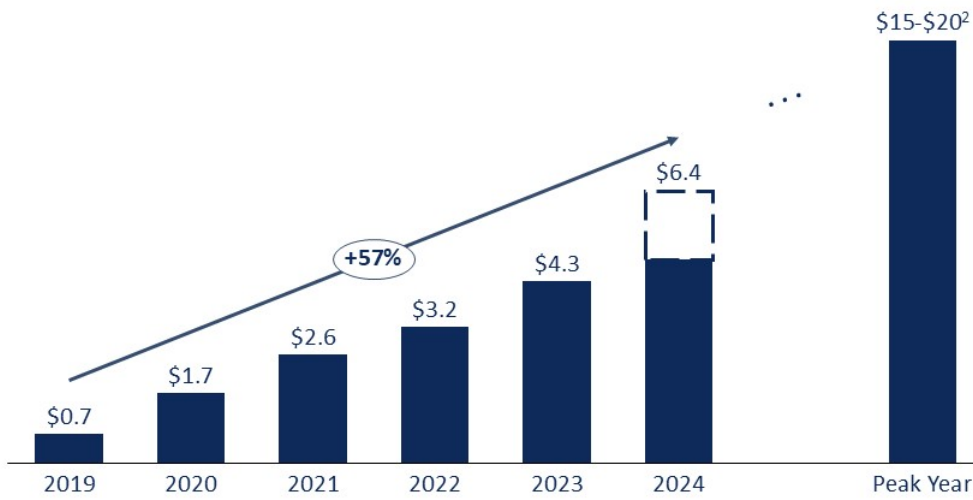
(acoramidis) 356 mg tablet



ATTR is a multi-billion-dollar market primed for continued expansion

Global annual ATTR market sales¹

\$B

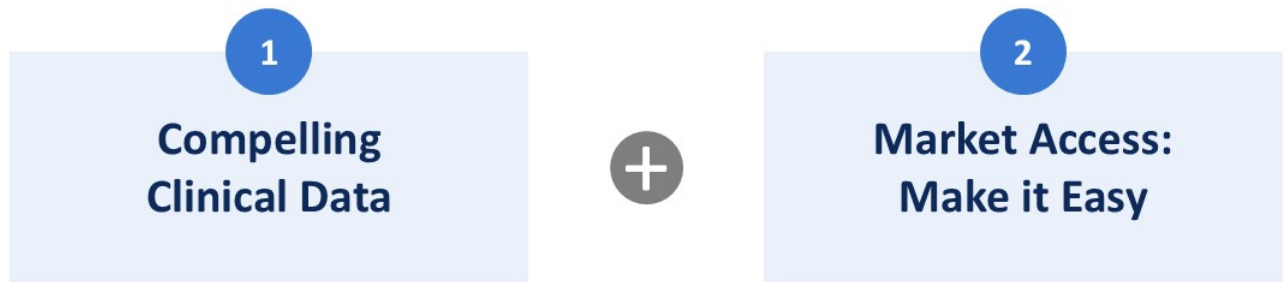


Market growth drivers include:

- With **more sponsors**, there is expanding disease awareness
- Increased global adoption of **non-invasive diagnostic tools**

¹ATTR market includes all approved drugs for ATTR-PN and ATTR-CM. 2024 sales annualized as of Q3 2024.
²Consensus estimates of \$15B-\$20B ATTR market.

The way we win





➤ **3 Months**

The most **rapid benefit** seen in any Phase 3 study of ATTR-CM to date

➤ **42% Reduction**

In **composite** of all-cause mortality and recurrent cardiovascular-related hospitalization events at Month 30

➤ **50% Reduction**

In the cumulative frequency of **cardiovascular-related hospitalization** events at Month 30

1 Data: ATTRuby achieved near-complete stabilization of TTR

Increase in serum TTR levels reflect *in vivo* stabilization

“Change from baseline in serum TTR levels were higher in participants receiving acoramidis only than those receiving placebo+tafamidis at Month 30...”¹

Increase in serum TTR levels observed with acoramidis treatment in patients with transthyretin amyloid cardiomyopathy (ATTR-CM): insights from ATTRIBUTE-CM and its open-label extension

1. Maurer, M. et al., Eur. Heart Jour., 2024;45(Suppl 1). 2. Nelson, L. et al., Amyloid, 2021;28(1):24-29. 3. Penchala, S. et al., PNAS, 2013;110(24):9992-97.

+ Evidence of potency on TTR stabilization

“AG10 [acoramidis] is 4 times more potent than tafamidis at a fixed plasma concentration (e.g., 10 μM).”²

Blinded potency comparison of transthyretin kinetic stabilisers by subunit exchange in human plasma

Luke T. Nelson^a, Ryan J. Paxman^a, Jin Xu^a, Bill Webb^b, Evan T. Powers^a and Jeffery W. Kelly^{a,c}

+ Differential TTR selectivity

“It is clear that AG10 [acoramidis] binds more selectively to TTR in serum than tafamidis.”³

AG10 inhibits amyloidogenesis and cellular toxicity of the familial amyloid cardiomyopathy-associated V122I transthyretin

Sravan C. Penchala^{a,1}, Stephen Connelly^{b,c,1}, Yu Wang^{a,1}, Miki S. Park^a, Lei Zhao^c, Aleksandra Baranczak^c, Irit Rappley^c, Hannes Vogel^d, Michaela Liedtke^d, Ronald M. Witteles^c, Evan T. Powers^c, Natàlia Reixach^c, William K. Chan^a, Ian A. Wilson^b, Jeffery W. Kelly^c, Isabella A. Graef^{d,2}, Mamoun M. Alhamadshah^{a,2}

+ Preclinical signals of superior potency & selectivity

“We carried out the subunit exchange in human plasma to address the relative selectivity of AG10 [acoramidis] vs. tafamidis...it is obvious that tafamidis is inferior to AG10 [acoramidis], but nothing like the degree you claim it is.”

–Prof. Jeffery Kelly (inventor of tafamidis) in email correspondence with Dr. Isabella Graef, February 27, 2013.

¹Maurer, M. et al., Eur. Heart Jour., 2024;45(Suppl 1). ²Nelson, L. et al., Amyloid, 2021;28(1):24-29. ³Penchala, S. et al., PNAS, 2013;110(24):9992-97.

2 Make it easy: simplified, differentiated, generous access programs



Free Trial

28-day supply for patients new to Attruby
(newly diagnosed or switching from another therapy)

Distribution Network

Independent and rare/orphan disease-focused Specialty Pharmacies (SPs) where Institutional SPs can keep prescription

Field Access Teams

Dedicated Attruby Field Reimbursement Managers and Patient Access Liaisons

Patient Assistance Programs

Easily accessible Copay and Patient Assistance Program

Commitment to clinical trial patients

US patients who participated in the acoramidis clinical trials may receive Attruby at no cost for the duration of their medically indicated treatment

Performance to date indicates strong commercial momentum



430

Attributed scripts
written to date



248

Unique
prescribing HCPs



77%

Medicare lives in equal
formulary position to tafamidis

Positive feedback all around on Market Access:

Pricing:

“ You priced responsibly and have a good label; we will have you at parity with tafamidis. Also, I have never seen a company offer free drug to their trial patients. Shows commitment.” - **Payer**

“ You did the right thing by being less than tafamidis... both your label and price will play a role here.” - **Payer**

Limited Distribution Network:

“ I want to reemphasize our gratitude for LDN inclusion. We objectively see key metrics in the patient journey improve as a result of dispensing access...Eager to leverage our resources to impact patient care in a positive way.” -**Manager of Health System SP**

“ You are a smart company to set up your distribution this way. I wish all companies would do this!” - **Director of Health System SP**

ForgingBridges:

“BridgeBio’s thoughtful resources and the structure of the network...aligns with the health system’s mission of putting patients first. - **Pharmacist**”

“Holy Smokes!? I wish all manufacturers would provide this information. Do you have any idea how much time this is going to save our team? - **Pharmacist**”

“Given the combination of strong efficacy and safety data, alongside BridgeBio’s very health system friendly approach towards access for this product (in comparison to the many challenges we have had with Pfizer...), we plan to begin preferring Attruby. - **AD Specialty Clinical Program Development**”

“It takes less than 5 minutes- the easiest enrollment process of any drug we use. - **RN Heart Failure Coordinator**”

Our Attruby team has experienced industry leaders who have built and launched blockbuster drugs



Matt Outten

Chief Commercial Officer

- Broad commercial leadership expertise with success across multiple competitive markets
- Led \$5B+ portfolio, 12 FDA approvals spanning 6 disease states and 7 indications (IMBRUVICA, Pharmacylics)
- Commercial lead on \$21B pharma M&A deal



Julie Everett

Chief Business Officer

- Successfully led cross-functional teams through multiple rare disease launches, including VOXZOGO and PALYNZIQ (BioMarin)
- Led commercial strategy/execution across ~\$1B portfolio
- ~Decade of strategy consulting leadership focused on launch excellence and lifecycle maximization (Trinity)



John Whang

Chief Medical Affairs Officer

- Orchestrated multiple successful launches with pioneering therapies in competitive segments – STELARA (Janssen), REPATHA (Amgen), and CAMZYOS (BMS)
- 8+ launches as strategy consultant (McKinsey)
- Demonstrated strategic innovation (Heartline Study – J&J / Apple collab) and consistently built outstanding organizations



Ana Merz

VP, Sales

Launched IMBRUVICA (\$5B+, 12 FDA approvals, 6 disease states, 7 indications in 10 years) and EPKINLY (3L+ DLBCL)



Sean Doherty

SVP, Marketing

Broad global sales and marketing launch experience including in rare, infectious, and auto-immune diseases



Scott Collins

SVP, Market Access

Extensive market access experience with consistent coverage across rare disease and oncology, leading large field-based access teams



Hudson Boyer

VP, Commercial Analytics & Ops

Launches in rare disease, hematology, and immunology; strategy consulting and equity analyst background



Liz Arnold

Head of Commercial Strategy

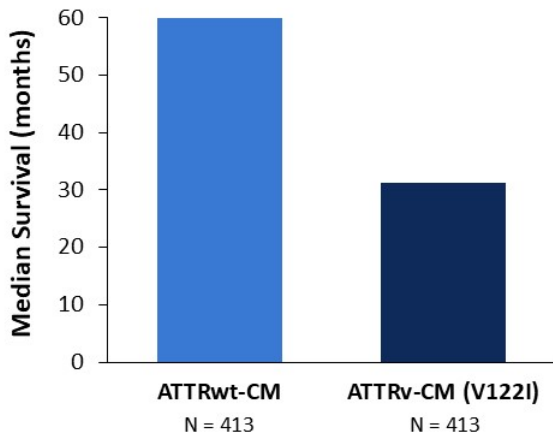
Multiple global launches, expertise in strategy, consulting, and marketing across rare disease, hematology, and OTC

Our BridgeBio team is committed to **providing industry-leading access and white glove service** for all parties looking to bring Attruby to patients with ATTR-CM.

Looking Ahead: Attruby Delivers Outstanding Results in Patients with Poor Prognosis

Natural History

V122I ATTRv-CM has an aggressive phenotype and poor prognosis¹



ATTRibute-CM mITT Population

Statistically significant benefit on composite ACM or first CVH in ATTRv-CM participants vs. placebo²

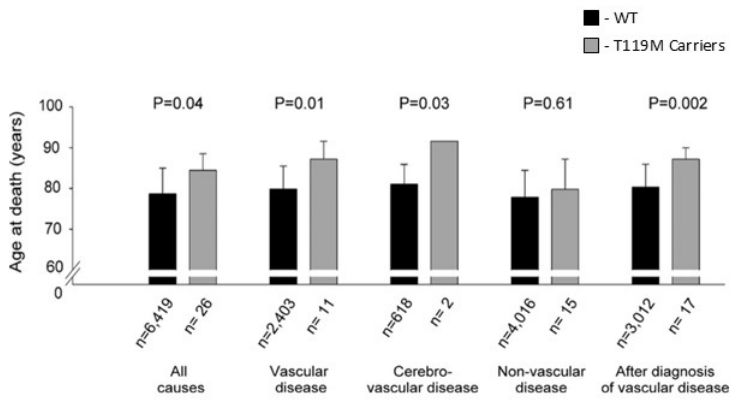
	N (%)	Hazard Ratio (95% CI)	p value
Overall Population	611 (100%)	0.65 (0.50-0.83)	0.0008
ATTRv-CM	59 (9.7%)	0.41 (0.21-0.81)	0.0109



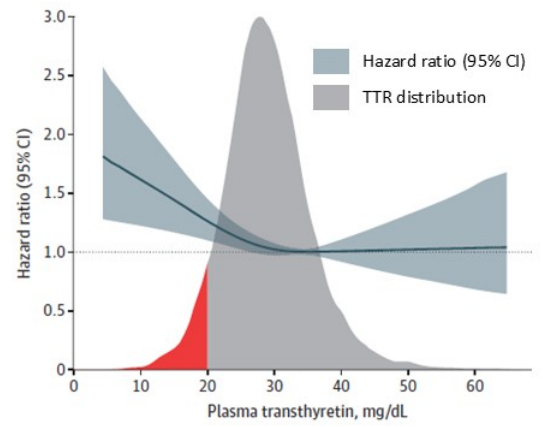
¹Razvi, Y. et al., Eur Jour Heart Fail., 2024;26(2):383-393. ²BridgeBio Data on File. Overall population and ATTRv-CM data vs. placebo.

Additional data: Elevated TTR levels are associated with improved survival

Genetic stabilization of transthyretin associated with improved health outcomes (N=69K individuals)¹



Higher TTR concentration associated with greater life expectancy (N≈102K individuals)²



¹Hornstrup, L. et al. *Arterioscler Thromb Vasc Biol*, 2013;33:1441-47. ²Figure adapted from Christoffersen, M. et al., *JAMA Cardiology*, 2024.

Infigratinib



Infigratinib is an oral best-in-class FGFR3 inhibitor that targets achondroplasia and hypochondroplasia at their source

Impact Opportunity

55,000+ People

In the US/EU with **achondroplasia (ACH)**, the most common form of disproportionate short stature.

55,000+ People

In the US/EU with **hypochondroplasia (HCH)**, another form of disproportionate short stature.

Design Criteria



Target FGFR3-driven skeletal dysplasias **at their source**



Demonstrate safety with **low dosing**



Convenient oral ROA to reduce patient burden

Phase 2 Data



Annualized height velocity
(To the sponsor's knowledge, largest effect reported to date)



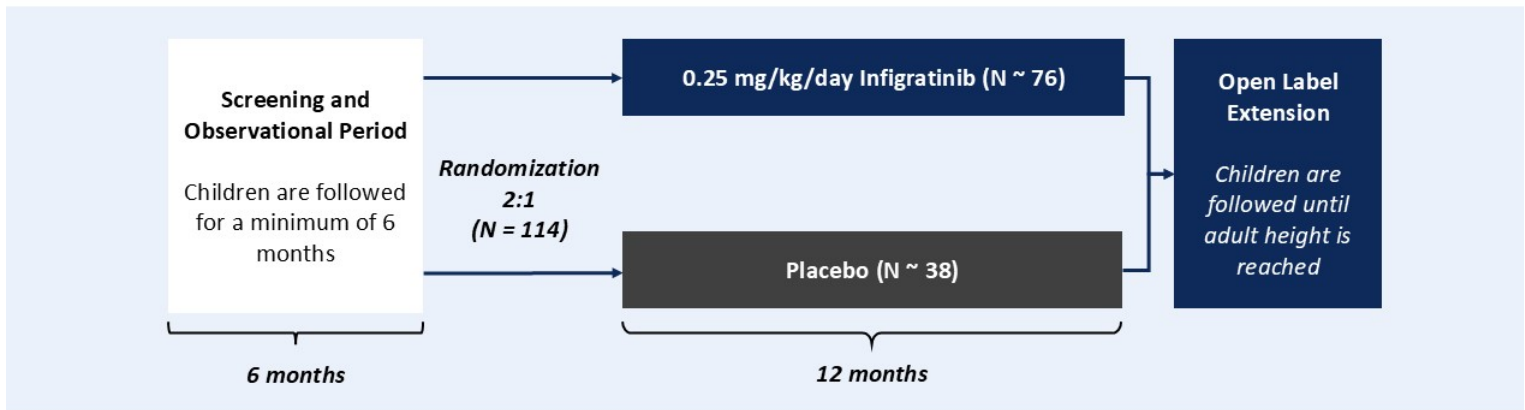
Proportionality
(To the sponsor's knowledge, largest effect reported to date)



Well-tolerated

\$4B+ Market Opportunity

We have fully enrolled a Phase 3 study (PROPEL 3) of Infigratinib in Achondroplasia with LPLV in 2H 2025



Primary Endpoint:

- Change from baseline in annualized height velocity at Wk 52

Key Secondary Endpoints:

- Change from baseline in height z-score
- Change from baseline in upper to lower body segment ratio

**Phase 3 Trial Fully Enrolled;
LPLV Expected 2H 2025**

BBP-418



BBP-418 is a first-in-class, disease-modifying therapy positioned to be the first approval in any form of limb-girdle muscular dystrophy

Impact Opportunity

7,000+ Patients

In the US/EU with LGMD2I/R9, a **progressive neuromuscular disorder** that leads to loss of ambulation, cardiomyopathy, and respiratory dysfunction.

Design Criteria



Targets the LMGD2I/R9 at its **source** by adding substrate to restore glycosylation of α DG



Avoid safety concerns by using a synthesized version of an endogenous compound



Convenient oral ROA to reduce patient burden

Phase 2 Data



Glycosylated α DG
(Expected surrogate endpoint)



Serum creatine kinase



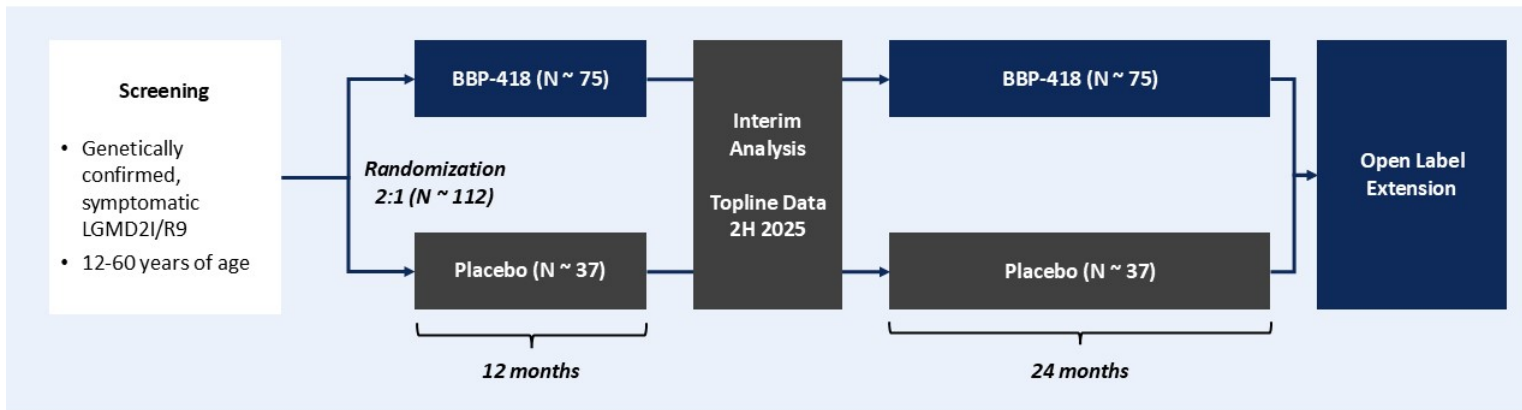
Stabilization of clinical measures



Well-tolerated

\$1B+ Market Opportunity + additional opportunity with other forms of LGMD

We have fully enrolled a Phase 3 study (FORTIFY) of BBP-418 in LGMD2I/R9 and expect topline interim analysis data readout in 2H 2025



Interim Endpoints:

- Glycosylated α DG (*primary*)
- Serum creatine kinase (CK)
- Trends in clinical measures

Final Analysis Endpoints:

- NSAD (*primary*)
- Ambulatory measures
 - 10MWT
 - 100MTT
- Pulmonary function: FVC
- Upper limb function: PUL 2.0
- QoL measures

**Phase 3 Trial Fully Enrolled;
LPLV & Topline Results
Expected 2H 2025**

Encaleret



Encaleret is a first-in-class, disease-modifying investigational therapy with potential to be the first approved intervention for ADH1

Impact Opportunity

25,000+ Patients

In the US/EU with **ADH1**, a genetic disease resulting in the disruption of Ca homeostasis.

200,000+ Patients

In the US/EU with **Post-Surgical Hypoparathyroidism (PSH)**, a dysregulation of Ca homeostasis caused by impaired parathyroid function.

Design Criteria



Target **ADH1 at its source** by desensitizing over-active Ca sensing receptors



Normalize PTH, serum Ca, and urine Ca levels



Convenient oral ROA to reduce patient burden

Phase 2 Data



Serum calcium to normal range



Urine calcium to normal range



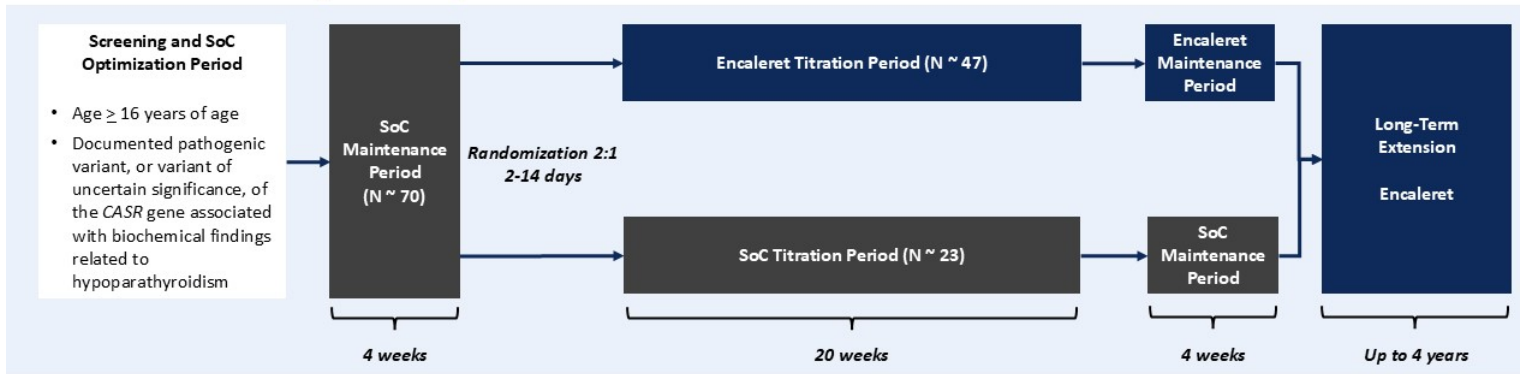
Parathyroid hormone to normal range



Well-tolerated

\$2B+ Market Opportunity

We have fully enrolled the Phase 3 study (CALIBRATE) of encaleret in ADH1 and expect topline results in 2H 2025



Primary Endpoint:

- Proportion of participants achieving:
 - Blood Ca within the target range **AND**
 - 24-hour urine Ca within the reference range

Select Secondary Endpoints:

- Blood iPTH, 1,25-(OH)₂ Vitamin D, magnesium, and phosphate
- Urine magnesium and phosphate
- Bone turnover markers
- Renal ultrasound and renal function
- ER/urgent care visits and/or hospitalizations
- Quality of life (SF-36)

**Phase 3 Trial Fully Enrolled;
LPLV & Topline Results
Expected 2H 2025**

SoC = standard of care; a combination of oral activated Vitamin D and/or calcium supplements. CASR = calcium-sensing receptor gene.

BBP-812
Canavan Disease



BBP-812 has the potential to change the disease trajectory for children with Canavan, a severe, ultra-rare, autosomal recessive leukodystrophy

*“One patient in particular is **sitting independently and taking steps and walking**, and that is certainly something I’ve never seen with Canavan disease.”*

– Dr. Florian Eichler, Mass General Hospital for Children

*“[The participant] was **like a new child, reactive, holding hands, and clapping.**”*

– Dr Annette Bley, University Medical Center Hamburg

*“Lots of improvement at 3-month visit, **emotional presence, visual tracking, reaching for things**”*

– Dr. Alexander Fay, UCSF Benioff Children's Hospital



Preliminary Data

- ↓ Urine NAA levels
- ↓ CNS NAA levels
- ↑ Gross Motor Skills
- ↑ Developmental Skills
- ✓ Safety data consistent with other AAV9 GTx's

BridgeBio
Ecosystem Highlights



BBOT has progressed two potentially first-in-class molecules into the clinic with a third expected in 1H 2025

Program	Mechanism of Action	Status
BBO-8520 KRAS ^{G12C} ON / OFF	<ul style="list-style-type: none">• First direct inhibitor of KRAS^{G12C} ON• Inhibits both KRAS^{G12C} GTP (active) and GDP (inactive) states• Differentiates from KRAS^{G12C} GDP (inactive)-only inhibitors	Enrolling
BBO-10203 RAS:PI3K α Breaker	<ul style="list-style-type: none">• Blocks specific interaction between RAS and PI3Kα• RAS driver agnostic (KRAS, HRAS and NRAS)• Selectively blocks PI3K / AKT effector signaling in the tumor• No hyperglycemia / hyperinsulinemia	Enrolling
BBO-11818 PanKRAS ON / OFF	<ul style="list-style-type: none">• Direct inhibitor of KRAS^{G12X} ON• Potent panKRAS inhibitor• Directly binds mutant KRAS	IND exp. Q1 2025

The GondolaBio pipeline features a diverse set of programs across TAs

Indication	Discovery	Lead Optimization	IND Enabling	Phase 1	Est. Patient Pop. (US + EU)
Erythropoietic Protoporphyrria (EPP)					20k
Galactosemia					10k
Alpha-1 Antitrypsin Deficiency (AATD)					200k
Neurofibromatosis Type 1 (NF1)					200k
Hereditary Pancreatitis					30k
Fibrous Dysplasia					50k
Congenital Glycosylation Disorder Type Ia (PMM2-CDG)					5k
Autosomal Dominant Polycystic Kidney Disease (ADPKD)					300k
Tuberous Sclerosis Complex 1/2 (TSC)					65k
Genetic Epilepsy Driven by SynGAP1 Mutations					10k

We are well-financed and expect to hit numerous milestones in 2025

1H 2025

- Acoramidis: EU approval
- Acoramidis: Japan approval
- Q1 Earnings Call

2H 2025

- Q2 and Q3 Earnings Calls
- Encaleret: LPLV, Topline
- BBP-418: LPLV, Topline
- Infigratinib: LPLV



bridgebio vision for 2030:



De-risked PYS
>\$8B



Lives impacted
>100k lives