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): August 4, 2022

AMICUS THERAPEUTICS, INC.

(Exact Name of Registrant as Specified in Its Charter)

Delaware
(State or Other Jurisdiction of Incorporation)

001-33497 (Commission File Number) 71-0869350 (I.R.S. Employer Identification No.)

3675 Market Street, Philadelphia, PA 19104 (Address of Principal Executive Offices, and Zip Code)

215-921-7600

Registrant's Telephone Number, Including Area Code

(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))
- $\begin{tabular}{ll} \hline \begin{tabular}{ll} \hline \end{tabular} \hline \end{tabular} \end{tab$

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 \$0.01	FOLD	NASDAQ

Indicate by check mark whether the registrant is an emerging growth company as defined in Rule 405 of the Securities Act of 1933 (17 CFR $\S 230.405$) or Rule 12b-2 of the Securities Exchange Act of 1934 (17 CFR $\S 240.12b-2$). Emerging growth company \square

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. \Box

Item 2.02 Results of Operations and Financial Condition

On August 4, 2022, Amicus Therapeutics, Inc. (the "Company") issued a press release announcing its financial results for the fiscal quarter ended June 30, 2022. A copy of this press release is attached hereto as Exhibit 99.1. The Company will host a conference call and webcast on August 4, 2022 to discuss its second quarter results of operations. A copy of the conference call presentation materials is attached hereto as Exhibit 99.2. Both exhibits are incorporated herein by reference.

In accordance with General Instruction B.2. of Form 8-K, the information in this Current Report on Form 8-K and the Exhibits shall not be deemed "filed" for purposes of Section 18 of the Securities Exchange Act of 1934, as amended (the "Exchange Act"), or otherwise subject to the liability of that section, nor shall it 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 filing.

Item 9.01 Financial Statements and Exhibits

(d) Exhibits:

Exhibit No.	Description
<u>99.1</u>	Press Release dated August 4, 2022
<u>99.2</u>	August 4, 2022 Conference Call Presentation Materials
104	Cover Page Interactive Data File (embedded within the Inline XBRL document)

Signature Page

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.

AMICUS THERAPEUTICS, INC.

By: /s/ Ellen S. Rosenberg

Name: Ellen S. Rosenberg

Title: Chief Legal Officer and Corporate Secretary

Date: August 4, 2022



Amicus Therapeutics Announces Second Quarter 2022 Financial Results

1H22 Galafold® Revenue of \$159.4M, reflecting 11% Sales Growth with Operational Growth of 18%, Partly Offset by Currency Headwinds of 7%

On-Track to Deliver Full-Year Double-Digit Revenue Growth of 15-20% at Constant Exchange Rates

Advancing U.S. and EU Regulatory Reviews and Launch Preparations for AT-GAA in Pompe Disease

3 Newly Issued U.S. Composition of Matter Patents for Galafold Add to Growing U.S. Patent Portfolio

Conference Call and Webcast Today at 8:30 a.m. ET

PHILADELPHIA, PA, Aug. 4, 2022 – Amicus Therapeutics (Nasdaq: FOLD), a patient-dedicated global biotechnology company focused on developing and commercializing novel medicines for rare diseases, today announced financial results for the quarter ended June 30, 2022.

Bradley Campbell, President and Chief Executive Officer of Amicus Therapeutics, Inc., stated, "Through the first half of the year and into the third quarter, we have gained great momentum towards achieving our key strategic priorities for 2022. We are pleased by the continued global uptake of Galafold and continued patient demand, which is driving strong operational growth in-line with our 2022 guidance. We are focused on gaining regulatory approvals of AT-GAA for people living with Pompe disease around the world. Importantly, we are poised for the anticipated successful launch of AT-GAA and continue to believe in the potential of this treatment regimen to become the new global standard of care in Pompe disease. These efforts, together with our careful management of expenses and the financial strength of our business, uniquely position Amicus to deliver sustainable value for shareholders while upholding our mission for people living with rare diseases."

Corporate Highlights

Global revenue for Galafold® (migalastat) in the first half of 2022 was \$159.4M, reflecting 11% growth with operational growth of 18% offset by currency headwinds of 7%. Second quarter sales of \$80.7 million represented a year-over-year increase of 13% at constant exchange rates (CER)¹. Second quarter reported revenue growth was 4% given significant currency headwinds of \$6.7 million, or 9%.

(in thousands)	1	Three Months Ended June 30,			Year over Year % Growth			Six Months E	Inde	d June 30,	Year over Year % Growth	
		2022		2021	As Reported	at CER ¹		2022		2021	As Reported	at CER ¹
Galafold Net Product Revenues	\$	80,731	\$	77,413	4%	13%	\$	159,446	\$	143,815	11%	18%

- Galafold U.S. intellectual property estate strengthened following the issuance of 17 new patents this year. The Galafold U.S. intellectual property portfolio now includes 44 orange book listed patents, 28 of which provide protection through at least 2038, including 3 newly issued composition of matter patents.
- AT-GAA regulatory reviews progressing and pre-launch activities underway. In the U.S., the Food and Drug Administration (FDA) extended the Prescription Drug User Fee Act (PDUFA) action dates to August 29, 2022, for the New Drug Application (NDA) and October 29, 2022, for the Biologic License Application (BLA), reflective of the two components of AT-GAA. The Company continues to expect the FDA to approve the applications together by the October 29, 2022, action date. In the EU, the Committee for Medicinal Products for Human Use (CHMP) opinion is expected in late 2022.
- Expanded access programs in place to meet the growing demand for AT-GAA across multiple countries. In France, the National Agency for the Medicines and Health Products Safety (ANSM) granted the first reimbursed access to AT-GAA under their compassionate access ("Accès Compassionnel") program. In the U.K., under the Early Access to Medicines Scheme (EAMS) multiple physicians have requested access across the leading Pompe centers in the country. Additional expanded access programs are in place in Germany and Japan with multiple Pompe patients participating in each.



- Amicus announces a program to explore next-generation pharmacological chaperones for Fabry disease through academic research collaboration agreement with the Spanish National Research Council (CSIC) and the University of Seville. This new collaboration will search for innovative glycomimetics with optimal activity and pharmacokinetic properties. Lead compounds will be selected based on their potential for greater potency, expanded number of amenable mutations, and optimal dosing.
- · Company leadership transition complete. As of August 1, 2022, Bradley Campbell transitioned to the role of President and Chief Executive Officer of Amicus. John F. Crowley has transitioned to the role of Executive Chairman of Amicus for a two-year term, after which he is expected to continue as the non-executive Chairman of the Board.
- On track to achieve non-GAAP profitability² in 2023. Through careful management of expenses, based on current operating models, the Company is on the path to achieve profitability in 2023, as it executes on the global expansion of Galafold and prepares for the global launch of AT-GAA.

Second Quarter 2022 Financial Results

- Total revenue in the second quarter 2022 was \$80.7 million, a year-over-year increase of 4% from total revenue of \$77.4 million in the second quarter of 2021. On a constant currency basis, second quarter 2022 total revenue growth was 13%. Reported revenue was offset by a negative currency impact of \$6.7 million, or 9%.
- · Cash, cash equivalents, and marketable securities totaled \$386.8 million at June 30, 2022, compared to \$482.5 million at December 31, 2021.
- · Total GAAP operating expenses of \$133.1 million for the second quarter 2022 increased as compared to \$107.9 million for the second quarter 2021.
- Total non-GAAP operating expenses of \$119.2 million for the second quarter of 2022 increased as compared to \$93.5 million in the second quarter of 2021, reflecting non-recurring expenses related to the reprioritization of the gene therapy portfolio.³
- Net loss was \$62.2 million, or \$0.21 per share, compared to a net loss of \$51.2 million, or \$0.19 per share, for the second quarter 2021.

2022 Financial Guidance

- For the full-year 2022, the Company anticipates total Galafold revenue of \$350 million to \$365 million at constant exchange rates¹. Double-digit revenue growth between 15 and 20% at CER¹ in 2022 is expected to be driven by continued underlying demand from both switch and treatment-naïve patients, geographic expansion, the continued diagnosis of new Fabry patients and commercial execution across all major markets, including the U.S., EU, U.K., and Japan. Applying average July 2022 exchange rates, the negative currency impact on full-year 2022 Galafold reported sales would be approximately 9%.
- Non-GAAP operating expense guidance for the full-year 2022 is \$470 million to \$485 million, driven by continued investment in the global Galafold launch, AT-GAA clinical studies and pre-launch activities, in addition to certain non-recurring costs for manufacturing to support the global launch of AT-GAA and committed obligations for the gene therapy portfolio. In 2023, Amicus expects non-GAAP operating expense at a level below 2021.

Anticipated 2022 Milestones by Program

Galafold (migalastat) Oral Precision Medicine for Fabry Disease

- · Sustain double-digit revenue growth in 2022 of \$350 million to \$365 million at CER
- · Continue geographic expansion
- · Registry and other Phase 4 studies ongoing

AT-GAA for Pompe Disease

· U.S. Prescription Drug User Fee Act (PDUFA) action date of August 29, 2022, for the NDA and October 29, 2022, for the BLA



- EU Committee for Medicinal Products for Human Use (CHMP) opinion expected in late 2022
- · Continue to broaden expanded access plans in the U.K., Germany, France, Japan, and other countries
- · Ongoing supportive studies, including pediatric and extension studies
- ¹ In order to illustrate underlying performance, Amicus discusses its results in terms of constant exchange rate (CER) growth. This represents growth calculated as if the exchange rates had remained unchanged from those used in the comparative period. Full-year 2022 Galafold revenue guidance utilizes the average actual exchange rates for 2021.
- ² Based on projections of Amicus non-GAAP Net Income under current operating plans, which includes successful AT-GAA regulatory approvals and continued Galafold growth. We define non-GAAP Net Income as GAAP Net Income excluding the impact of share-based compensation expense, changes in fair value of contingent consideration, loss on impairment of assets, depreciation and amortization, acquisition related income (expense), loss on extinguishment of debt, loss on impairment of assets, restructuring charges and income taxes.
- ³ Full reconciliation of GAAP results to the Company's non-GAAP adjusted measures for all reporting periods appear in the tables to this press release.
- ⁴ A reconciliation of the differences between the non-GAAP expectation and the corresponding GAAP measure is not available without unreasonable effort due to high variability, complexity, and low visibility as to the items that would be excluded from the GAAP measure.

Conference Call and Webcast

Amicus Therapeutics will host a conference call and audio webcast today, August 4, 2022, at 8:30 a.m. ET to discuss the second quarter 2022 financial results and corporate updates. Interested participants and investors may access the conference call by dialing 1-833-634-2601 (U.S. Toll Free) or 1-412-902-4113 (International).

A live audio webcast and related presentation materials can also be accessed via the Investors section of the Amicus Therapeutics corporate website at ir.amicusrx.com. Web participants are encouraged to register on the website 15 minutes prior to the start of the call. A replay of the call will be available for seven days beginning two hours after the conclusion of the event. Access numbers for this replay are 1-877-344-7529 (U.S. Toll Free), 1-855-669-9658 (Canada Toll Free), and 1-412-317-0088 (International); Access Code: 3033146.

About Galafold

Galafold[®] (migalastat) 123 mg capsules is an oral pharmacological chaperone of alpha-Galactosidase A (alpha-Gal A) for the treatment of Fabry disease in adults who have amenable galactosidase alpha gene (GLA) variants. In these patients, Galafold works by stabilizing the body's own dysfunctional enzyme so that it can clear the accumulation of disease substrate. Globally, Amicus Therapeutics estimates that approximately 35 to 50 percent of Fabry patients may have amenable GLA variants, though amenability rates within this range vary by geography. Galafold is approved in more than 40 countries around the world, including the U.S., EU, U.K., and Japan.

U.S. INDICATIONS AND USAGE

Galafold is indicated for the treatment of adults with a confirmed diagnosis of Fabry disease and an amenable galactosidase alpha gene (GLA) variant based on in vitro assay data.

This indication is approved under accelerated approval based on reduction in kidney interstitial capillary cell globotriaosylceramide (KIC GL-3) substrate. Continued approval for this indication may be contingent upon verification and description of clinical benefit in confirmatory trials.

U.S. IMPORTANT SAFETY INFORMATION

ADVERSE REACTIONS

The most common adverse reactions reported with Galafold (≥10%) were headache, nasopharyngitis, urinary tract infection, nausea and pyrexia.

USE IN SPECIFIC POPULATIONS

There is insufficient clinical data on Galafold use in pregnant women to inform a drug-associated risk for major birth defects and miscarriage. Advise women of the potential risk to a fetus.

It is not known if Galafold is present in human milk. Therefore, the developmental and health benefits of breastfeeding should be considered along with the mother's clinical need for Galafold and any potential adverse effects on the breastfeed child from Galafold or from the underlying maternal condition.

Galafold is not recommended for use in patients with severe renal impairment or end-stage renal disease requiring dialysis.

The safety and effectiveness of Galafold have not been established in pediatric patients.

To report Suspected Adverse Reactions, contact Amicus Therapeutics at 1-877-4AMICUS or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

For additional information about Galafold, including the full U.S. Prescribing Information, please visit https://www.amicusrx.com/pi/Galafold.pdf.



EU Important Safety Information

Treatment with Galafold should be initiated and supervised by specialists experienced in the diagnosis and treatment of Fabry disease. Galafold is not recommended for use in patients with a nonamenable mutation.

- · Galafold is not intended for concomitant use with enzyme replacement therapy
- · Galafold is not recommended for use in patients with Fabry disease who have severe renal impairment (<30 mL/min/1.73 m2). The safety and efficacy of Galafold in children less than 12 years of age have not yet been established. No data are available.
- · No dosage adjustments are required in patients with hepatic impairment or in the elderly population.
- · There is very limited experience with the use of this medicine in pregnant women. If you are pregnant, think you may be pregnant, or are planning to have a baby, do not take this medicine until you have checked with your doctor, pharmacist, or nurse.
- · While taking Galafold, effective birth control should be used. It is not known whether Galafold is excreted in human milk
- · Contraindications to Galafold include hypersensitivity to the active substance or to any of the excipients listed in the PRESCRIBING INFORMATION.
- · Galafold 123 mg capsules are not for children (≥12 years) weighing less than 45 kg.
- · It is advised to periodically monitor renal function, echocardiographic parameters and biochemical markers (every 6 months) in patients initiated on Galafold or switched to Galafold.
- · OVERDOSE: General medical care is recommended in the case of Galafold overdose
- The most common adverse reaction reported was headache, which was experienced by approximately 10% of patients who received Galafold. For a complete list of adverse reactions, please review the SUMMARY OF PRODUCT CHARACTERISTICS.
- · Call your doctor for medical advice about side effects

For further important safety information for Galafold, including posology and method of administration, special warnings, drug interactions and adverse drug reactions, please see the European SmPC for Galafold available from the EMA website at www.ema.europa.eu.

About Amicus Therapeutics

Amicus Therapeutics (Nasdaq: FOLD) is a global, patient-dedicated biotechnology company focused on discovering, developing and delivering novel high-quality medicines for people living with rare metabolic diseases. With extraordinary patient focus, Amicus Therapeutics is committed to advancing and expanding a robust pipeline of cutting-edge, first- or best-in-class medicines for rare metabolic diseases. For more information please visit the company's website at www.amicusrx.com. and follow on Twitter and LinkedIn.

Non-GAAP Financial Measures

In addition to financial information prepared in accordance with U.S. GAAP, this press release also contains adjusted financial measures that we believe provide investors and management with supplemental information relating to operating performance and trends that facilitate comparisons between periods and with respect to projected information. These adjusted financial measures are non-GAAP measures and should be considered in addition to, but not as a substitute for, the information prepared in accordance with U.S. GAAP. We typically exclude certain GAAP items that management does not believe affect our basic operations and that do not meet the GAAP definition of unusual or non-recurring items. Other companies may define these measures in different way. Full reconcilations of GAAP results to the comparable non-GAAP measures for the reported periods appear in the financial tables section of this press release. When we provide our expectation for non-GAAP operating expenses on a forward-looking basis, a reconciliation of the differences between the non-GAAP expectation and the corresponding GAAP measure generally is not available without unreasonable effort due to potentially high variability, complexity and low visibility as to the items that would be excluded from the GAAP measure in the relevant future period, such as unusual gains or losses. The variability of the excluded items may have a significant, and potentially unpredictable, impact on our future GAAP results.



Forward-Looking Statements

This press release contains "forward-looking statements" within the meaning of the Private Securities Litigation Reform Act of 1995 relating to preclinical studies and clinical trials, the prospects and timing of the potential regulatory approval of our product candidates, commercialization plans, manufacturing and supply plans, financing plans, and the projected revenues, expenses, cash position, and future profitability for the Company. The inclusion of forward-looking statements should not be regarded as a representation by us that any of our plans will be achieved. Any or all of the forward-looking statements in this press release may turn out to be wrong and can be affected by inaccurate assumptions we might make or by known or unknown risks and uncertainties. For example, with respect to statements regarding the goals, progress, timing, and outcomes of discussions with regulatory authorities, and in particular the potential goals, progress, timing, and results of preclinical studies and clinical trials, including as they are impacted by COVID-19 pandemic is inherently unknown and cannot be predicted with confidence and may cause actual results and performance to differ materially from the statements in this release, including without limitation, because of the impact on general political and economic conditions, including as a result of efforts by governmental authorities to mitigate COVID-19, such as travel bans, shelter in place orders and third-party business closures and resource allocations, manufacturing and supply chain disruptions and limitations on patient access to commercial or clinical product. In addition to the impact of the COVID-19 pandemic, actual results may differ materially from those set forth in this release due to the risks and uncertainties inherent in our business, including, without limitation: the potential that it may be difficult to erroll patients in our clinical trials; the potential that regulatory authorities, including the FDA, EMA, and PMDA, may not grant or may delay ap

CONTACTS:

Investors:

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Media:

Amicus Therapeutics Diana Moore Head of Global Corporate Communications dmoore@amicusrx.com (609) 662-5079

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TABLE 1

Amicus Therapeutics, Inc. Consolidated Statements of Operations (Unaudited) (in thousands, except share and per share amounts)

	Three Months Ended June 30,			Six Months Ended June 30,			
		2022		2021	 2022		2021
Net product sales	\$	80,731	\$	77,413	\$ 159,446	\$	143,815
Cost of goods sold		8,197		8,380	15,779		14,919
Gross profit		72,534		69,033	143,667		128,896
Operating expenses:							
Research and development		78,319		63,003	159,836		127,120
Selling, general, and administrative		53,379		42,276	111,495		89,002
Changes in fair value of contingent consideration payable		115		1,021	(1,073)		1,492
Loss on impairment of assets		_		_	6,616		_
Depreciation and amortization		1,334		1,567	2,745		3,171
Total operating expenses		133,147		107,867	279,619		220,785
Loss from operations		(60,613)		(38,834)	(135,952)		(91,889)
Other (expense) income:							
Interest income		356		50	489		215
Interest expense		(8,257)		(8,150)	(16,404)		(16,142)
Other income		7,268		234	9,170		(2,966)
Loss before income tax		(61,246)		(46,700)	(142,697)		(110,782)
Income tax expense		(911)		(4,525)	(4,720)		(6,107)
Net loss attributable to common stockholders	\$	(62,157)	\$	(51,225)	\$ (147,417)	\$	(116,889)
Net loss attributable to common stockholders per common share — basic and diluted	\$	(0.21)	\$	(0.19)	\$ (0.51)	\$	(0.44)
Weighted-average common shares outstanding — basic and diluted		291,970,562		266,398,516	288,646,587		265,384,865



TABLE 2

Amicus Therapeutics, Inc. Consolidated Balance Sheets (Unaudited) (in thousands, except share and per share amounts)

		June 30, 2022		ecember 31, 2021
Assets				
Current assets:				
Cash and cash equivalents	\$	235,639	\$	245,197
Investments in marketable securities		151,202		237,299
Accounts receivable		52,556		52,672
Inventories		20,879		26,818
Prepaid expenses and other current assets		37,367		34,848
Total current assets		497,643		596,834
Operating lease right-of-use assets, net		30,447		20,586
Property and equipment, less accumulated depreciation of \$22,188 and \$19,882 at June 30, 2022 and December 31, 2021, respectively		33,657		42,496
In-process research & development		23,000		23,000
Goodwill		197,797		197,797
Other non-current assets		18,045		24,427
Total Assets	\$	800,589	\$	905,140
Liabilities and Stockholders' Equity				
Current liabilities:				
Accounts payable	\$	23,113	\$	21,513
Accrued expenses and other current liabilities		114,703		98,153
Contingent consideration payable		19,266		18,900
Operating lease liabilities		7,543		7,409
Total current liabilities		164,625		145,975
Long-term debt		390,652		389,357
Operating lease liabilities		52,844		43,363
Deferred reimbursements		5,906		5,906
Deferred income taxes		4,930		4,930
Other non-current liabilities		8,207		8,240
Total liabilities	_	627,164		597,771
Commitments and contingencies		•		
Stockholders' equity:				
Common stock, \$0.01 par value, 500,000,000 shares authorized, 280,456,667 and 278,912,800 shares issued and outstanding at June 30, 2022 and December 31, 2021,				
respectively		2,811		2,808
Additional paid-in capital		2,631,110		2,595,419
Accumulated other comprehensive (loss) gain:				
Foreign currency translation adjustment		(16,603)		5,251
Unrealized loss on available-for-sale securities		(637)		(270)
Warrants		83		83
Accumulated deficit		(2,443,339)		(2,295,922)
Total stockholders' equity		173,425		307,369
Total Liabilities and Stockholders' Equity	\$	800,589	\$	905,140
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TABLE 3

Amicus Therapeutics, Inc. Reconciliation of Non-GAAP Financial Measures (in thousands)

		Three Months Ended June 30,				Six Months Ended June 30,			
	2	022		2021		2022		2021	
Total operating expenses - as reported GAAP	\$	133,147	\$	107,867	\$	279,619	\$	220,785	
Research and development:									
Share-based compensation		4,379		3,152		13,744		9,457	
Selling, general and administrative:									
Share-based compensation		8,084		8,584		29,370		22,633	
Loss on impairment of assets		_		_		6,616		_	
Changes in fair value of contingent consideration payable		115		1,021		(1,073)		1,492	
Depreciation and amortization		1,334		1,567		2,745		3,171	
Total operating expense adjustments to reported GAAP		13,912		14,324		51,402		36,753	
Total operating expenses - as adjusted	\$	119,235	\$	93,543	\$	228,417	\$	184,032	
	8								



2Q22 Financial Results Conference Call & Webcast

At the Forefront of Therapies for Rare Diseases

August 4, 2022



Forward-Looking Statements

This presentation contains "forward-looking statements" within the meaning of the Private Securities Litigation Reform Act of 1995 relating to preclinical and clinical development of our product candidates, the timing and reporting of results from preclinical studies and clinical trials, the prospects and timing of the potential regulatory approval of our product candidate commercialization plans, manufacturing and supply plans, financing plans, and the projected revenues, expenses, cash position, and future profitability for the Company. The inclusion forward-looking statements should not be regarded as a representation by us that any of our plans will be achieved. Any or all of the forward-looking statements in this press release may to out to be wrong and can be affected by inaccurate assumptions we might make or by known or unknown risks and uncertainties. For example, with respect to statements regarding the go progress, timing, and outcomes of discussions with regulatory authorities, and in particular the potential goals, progress, timing, and results of preclinical studies and clinical trials, including they are impacted by COVID-19 related disruption, are based on current information. The potential impact on operations from the COVID-19 pandemic is inherently unknown and cannot predicted with confidence and may cause actual results and performance to differ materially from the statements in this release, including without limitation, because of the impact on gene political and economic conditions, including as a result of efforts by governmental authorities to mitigate COVID-19, such as travel bans, shelter in place orders and third-party business closu and resource allocations, manufacturing and supply chain disruptions and limitations on patient access to commercial or clinical product. In addition to the impact of the COVID-19 pandem actual results may differ materially from those set forth in this release due to the risks and uncertainties inherent in our business, including, without limitation: the

Non-GAAP Financial Measures

In addition to financial information prepared in accordance with U.S. GAAP, this presentation also contains adjusted financial measures that we believe provide investors and management we supplemental information relating to operating performance and trends that facilitate comparisons between periods and with respect to projected information. These adjusted finance measures are non-GAAP measures and should be considered in addition to, but not as a substitute for, the information prepared in accordance with U.S. GAAP. We typically exclude cert GAAP items that management does not believe affect our basic operations and that do not meet the GAAP definition of unusual or non-recurring items. Other companies may define the measures in different ways. When we provide our expectation for non-GAAP operating expenses on a forward-looking basis, a reconciliation of the differences between the non-GA expectation and the corresponding GAAP measure generally is not available without unreasonable effort due to potentially high variability, complexity and low visibility as to the items to would be excluded from the GAAP measure in the relevant future period, such as unusual gains or losses. The variability of the excluded items may have a significant, and potentic unpredictable, impact on our future GAAP results.

10

A Rare Company

Patient-dedicated, Rare Disease Biotechnology Company with Sustained Double-digit Revenue Gro a Global Commercial Infrastructure, and Late-stage Development Capabilities



First Oral Precision Medicine for Fabry Disease

EMPLOYEES in 27 Countries

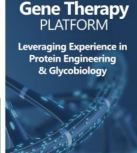
COMMERCIAL

ORGANIZATION

GLOBAL







\$350M-\$365M FY22 Global Galafold Revenue at CER World-class CLINICAL DEVELOPMENT Capabilities



Non-GAAP PROFITABILIT expected in 2023

GALAFOLD

AT-GAA

Cumulative \$2B

Peak Potential

\$386.8 Cash as of 6/30/2

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Positioned for Significant Value Growth

Focused on Execution and Driving Sustainable Double-digit Revenue Growth on Path to Profitability



Continue to bring Galafold® to as many patients as possible, sustain double-digit revenue growth



Successful launch of AT-GAA for people living with Pompe disease



Advance next-generation gene therapies in Fabry and Pompe diseases



Fully leverage global capabilities and infrastructure as a leader in rare diseases



Achieve non-GAAP profitability in 2023



2022 Strategic Priorities to Drive Value

- 1 Double-digit Galafold growth (15-20%) with revenue of \$350M to \$365M at CER¹
- 2 Secure FDA approval and positive CHMP opinion for AT-GAA
- 3 Initiate successful, rapid launch in U.S. for AT-GAA
- 4 Advance best-in-class, next-generation genetic medicines and capabilities
- Maintain strong financial position on path to profitability



Galafold® (migalastat) Continued Growth...

... building a leadership position in the treatment of Fabry disease



Galafold Success (as of June 30, 2022)

Building on Galafold's Success and Leveraging Leadership Position to Drive Continued Growth

Galafold is the first and only approved oral treatment option with a unique mechanism of action for Fabry patients with amenable variants



Galafold is indicated for adults with a confirmed diagnosis of Fabry Disease and an amenable variant. The most common adverse reactions reported with Galafold (≥10%) were headache, nasopharyngitis, urinary tract infection, nausea and pyrexia. For additional information about Galafold, including the full U.S. Prescribing information, please visit https://www.amicusrc.com/pi/Galafold.pdf For further important safety information for Galafold, including posology and method of administration, special warnings, drug interactions and adverse drug reactions, please see the European SmPC for Galafold available from the EMA website at www.ema.europa.eu.



7

Galafold Performance

1H22 Reported Revenue Growth of +10.9% to \$159.4M - Operational Growth of +17.8% at CEF



- Global 3-month net new patients trend greater than
 6-month and 12-month
- In the U.S., the month of June saw highest net new patients and PRFs since April 2021
- Global mix of switch (~55%) and previously untreated patients (~45%)
- Compliance and adherence over 90%+
- Expect non-linear quarterly growth to continue due t uneven ordering patterns



Galafold Success and FY22 Revenue Guidance

Galafold Momentum on Track to Achieve Full-year 2022 Revenue Guidance at CER



Reiterating FY22 Revenue Guidance of \$350M-\$365M at CER (between 15% and 20% growth at CER)

9 12022 Galafold revenue guidance utilizes the average actual rates for 2021



Galafold Growth Opportunity

\$1B Annual Sales Opportunity at Peak

> **Sustained** double-digit revenue growth:

1H operational revenue growth of +17.8% **Near-term** growth to \$500M driven by:

Continued penetration into existing markets

Expansion into new geographies

Broadening of labels

Long-term growth towards peak sales potential driven by:

Penetration of the diagnosed untreated population

Increase in newborn screening and diagnostic initiatives

Strong IP rights, including COM protection through 2038

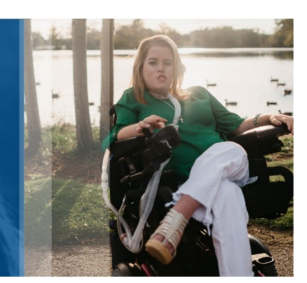


10 COM: Composition of Matter

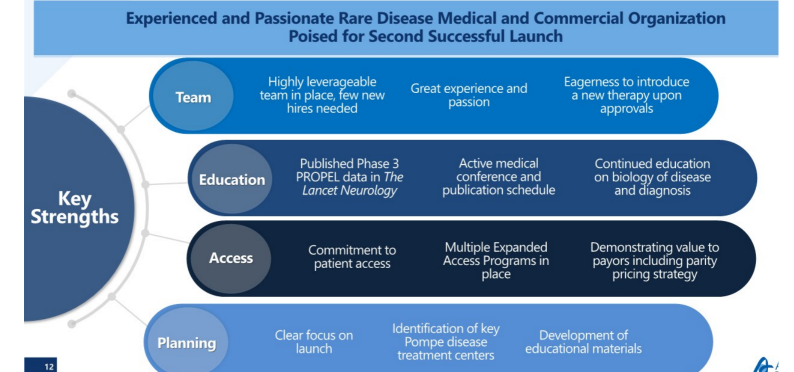


AT-GAA (cipaglucosidase alfa + miglustat)

... potential to establish a new standard of care for people living with Pompe disease



AT-GAA Launch Preparations



Pompe Disease Overview

Pompe is a Severe and Fatal Neuromuscular Disease Caused by the Deficiency of Lysosomal Enzyme G



Estimated incidence of ~1:28,000; newborn screening suggests significant underdiagnosis

Age of onset ranges from infancy to adulthood

Majority of patients on current standard of care decline after ~2 years Respiratory and cardiac failure are leading causes of morbidity and mortality

Deficiency of GAA leading to lysosomal glycogen accumulation and cellular dysfunction

Symptoms include muscle weakness, respiratory failure, and cardiomyopathy

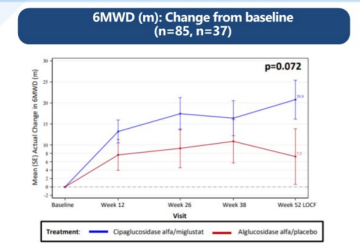
~\$1.2B+ global Pompe ERT sales¹

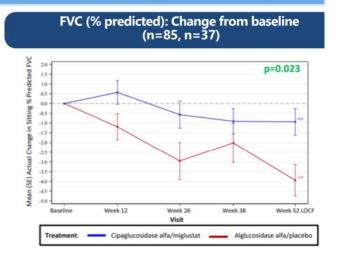


Phase 3 PROPEL Study Results

Overall Population (n=122*)

Primary and First Key Secondary Endpoint Showed Greater Improvement with AT-GAA vs. alglucosidase alfa in the Overall Population of ERT-Naïve and ERT-Experienced Patients





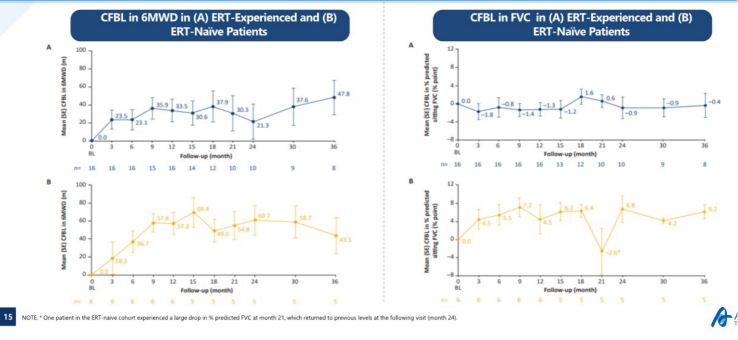
Results in ERT-Experienced Patients (n=92) Showed Meaningful Improvement for Both 6MWD (P=0.046) and FVC (P=0.006)

1

MMD=6-minute walk distance; EV=610 distance; DE estandard error. P values are nominal 2-sided; EVC data normally distributed and P value is from ANCOVA. 6MWD data not normally distributed and P value is from ANCOVA.

Long-Term Data from Phase 1/2 Clinical Study (ATB200-02)

Persistent and Durable Improvements in Motor and Respiratory Function and Reductions in Biomarkers of Muscle Damage and Disease Substrate Observed in Patients out to 36 Months



AT-GAA: Key Takeaways

Focused on Advancing AT-GAA to as Many Patients as Possible through Global Regulatory Pathways and Expanded Access Mechanisms

- Regulatory status update:
 - U.S. PDUFA date 2H2022¹
 - CHMP opinion late 2022
 - Planning for additional regulatory submissions
- Multiple expanded access mechanisms in place, including in the U.S., U.K., Germany, France, Japan, and others
 - First reimbursed access through the French compassionate access program
- 175+ people living with Pompe disease are now on AT-GAA across our clinical extension studies and expanded access programs
- Ongoing supportive studies:
 - Late-Onset Pompe Disease (LOPD) in children and adolescents aged 0 to <18
 - Infantile-Onset Pompe Disease (IOPD)





Amicus Pipeline

Streamlined Rare Disease Pipeline with Focus on Fabry Disease and Pompe Disease





Financial & Operational Strategy ... maintaining a strong financial outlook

Revenue Performance

Q2 Revenue Growth of +4.3% to \$80.7M resulting from Strong Operational Growth of +12.9% at CER Offset by Negative FX impact of -8.6%

Year-over-Year Sales Growth



- Significant currency exposure as 66% of Galafold revenue generated outside the U.
- Applying average July 2022 exchange rates the negative FX impact on full-year 2022 Galafold® reported sales would be approximately -9%, or ~\$26 million.



2Q Select Financial Results

2Q22 OpEx Increase Reflects Manufacturing Costs to Support AT-GAA Launch and Non-Recurring Expenses Related to the Reprioritization of the Gene Therapy Portfolio

	l 20 2022	
(in thousands, except per share data)	Jun. 30, 2022	Jun. 30, 2021
Product Revenue	\$80,731	\$77,413
Cost of Goods Sold	8,197	8,380
R&D Expense	78,319	63,003
SG&A Expense	53,379	42,276
Changes in Fair Value of Contingent Consideration	115	1,021
Depreciation and Amortization	1,334	1,567
Loss from Operations	(60,613)	(38,834)
Income Tax Expense	(911)	(4,525)
Net Loss	(62,157)	(51,225)
Net Loss Per Share	(0.21)	(0.19)



Financial Outlook and Path to Profitability

Clear Strategy to Build Our Business, Advance Our Portfolio, and Achieve Profitability



Sustain Galafold Revenue Growth

\$159.4M 1H2022 revenue, +17.8% YoY Operational Growth

2022 Galafold revenue guidance of \$350M-\$365M at CER, +15-20% YoY Growth



Secure Approvals of AT-GAA

Galafold and AT-GAA expected to drive strong double-digit growth long term



Deliver on Financial Goals

Focused on prudent expense management

2022 non-GAAP operating expense guidance of **\$470M-\$485M**

Achieve profitability¹ in 2023

¹Based on projections of Amicus non-GAAP Net Income under current operating plans, which includes successful AT-GAA regulatory approvals and continued Galafold growth. We define non-GAAP Net Income as GAAP Net Income excluding the impact of share-based compensation expense, changes in fair value of contingent consideration, depreciation and amortization, acquisition related income (expense), loss on extinguishment of debt, loss on impairment of assets, restructuring charges, and income taxes.







Amicus Therapeutics, Inc. Reconciliation of Non-GAAP Financial Measures (in thousands)

	Three Mon	ths Ended June 30,	Six Months Ended June 30,					
	2022	2021	2022	2021				
Total operating expenses - as reported GAAP	\$ 133,147	\$ 107,867	\$ 279,619	\$ 220,785				
Research and development:								
Share-based compensation	4,379	3,152	13,744	9,457				
Selling, general and administrative:								
Share-based compensation	8,084	8,584	29,370	22,633				
Loss on impairment of assets	<u></u>	_	6,616	_				
Changes in fair value of contingent	115	1,021	(1,073)	1,492				
consideration payable								
Depreciation and amortization	1,334	1,567	2,745	3,171				
Total operating expense adjustments to reported	13,912	14,324	51,402	36,753				
GAAP								
Total operating expenses - as adjusted	\$ 119,235	\$ 93,543	\$ 228,417	\$ 184,032				

