### 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): June 22, 2023

### Sarepta Therapeutics, Inc.

(Exact name of Registrant as Specified in Its Charter)

Delaware (State or Other Jurisdiction of Incorporation) 001-14895 (Commission File Number) 93-0797222 (IRS Employer Identification No.)

215 First Street Cambridge, Massachusetts (Address of Principal Executive Offices)

02142 (Zip Code)

Registrant's Telephone Number, Including Area Code: (617) 274-4000

(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)

Derecommencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))

D 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:

	Trading	
Title of each class	Symbol(s)	Name of each exchange on which registered
Common Stock, \$0.0001 par value per share	SRPT	Nasdag 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  $\Box$ 

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 7.01 Regulation FD Disclosure.

On June 22, 2023, Sarepta Therapeutics, Inc. (the "Company") conducted an investor webcast related to ELEVIDYS, its gene therapy product for the treatment of ambulatory pediatric patients aged 4 through 5 years with Duchenne muscular dystrophy ("DMD") with a confirmed mutation in the DMD gene. The Company disclosed in the webcast the United States wholesale acquisition cost (gross price) of ELEVIDYS at \$3.2 million. A copy of the Company's presentation is being furnished as Exhibit 99.1.

The information in this report, including Exhibit 99.1 attached hereto, is furnished pursuant to Item 7.01 and shall not be deemed "filed" for the 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. It may only be incorporated by reference in another filing under the Exchange Act or the Securities Act of 1933, as amended, if such subsequent filing specifically references the information furnished pursuant to Item 7.01 of this report.

#### Item 9.01. Financial Statements and Exhibits.

(d) Exhibits

Evhibit

Number	Description
99.1	Presentation dated June 22, 2023
104	The cover page from this Current Report on Form 8-K of Sarepta Therapeutics, Inc., formatted in Inline XBRL and included as Exhibit 101

### 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 thereunto duly authorized.

### Sarepta Therapeutics, Inc.

By: /s/ Douglas S. Ingram

Douglas S. Ingram President and Chief Executive Officer

Date: June 22, 2023

## So neither will we.

Exhibit 99.1

BENJAMIN Living with Duchenne muscular dystrophy

Doug Ingram President and CEO Sarepta Therapeutics, Inc. (NASDAQ: SRPT)

June 22, 2023



### **Forward-looking Statements**

This presentation contains "forward-looking statements." Any statements that are not statements of historical fact may be deemed to be forward-looking statements. Words such as "believe," "anticipate," "plan," "expect," "will," "may," "intend," "prepare," "look," "potential," "possible" and similar expressions are intended to identify forward-looking statements. These forward-looking statements include, without limitation, statements relating to our future operations, business plans, priorities, research and development programs; the potential benefits and risks of ELEVIDYS the expected demand for ELEVIDYS; our guidance for full-year 2023 net product revenue for our three approved oligonucleotide therapies of \$925 million dollars; our expectation that ELEVIDYS to halt the otherwise irreversible muscle damage caused by Duchenne; our belief that we have met our goal of ensuring that the costs to the healthcare system are less than the potential benefits of ELEVIDYS; the belief that nearly all infusions of ELEVIDYS will be subject to a statutory discount; our cost-effectiveness analysis; the potential pager budget impact; our understanding from the FDA that if the read out of EMBARK meets its objectives, the FDA intends to entertain a non-age restricted expansion of the label and that this will be done with maximal speed by the FDA; and expected plans and milestones, including our expectations that EMBARK, if the study meets its objectives, will act as our confirmatory status, if the trial is successful, our goal to expand the ELEVIDYS label to treat as much as 95% of the Duchenne population, if successful, and company to possible of ambulatory status, if the trial is successful, our goal to expand the ELEVIDYS label to treat as much as 95% of the Duchenne population, if successful, and commencing multiple trials to explore the clearance of pre-existing antibodies, in addition to ENVISION.

Actual results could materially differ from those stated or implied by these forward-looking statements as a result of such risks and uncertainties. Known risk factors include the following: the FDA may not approve a supplement to expand the approved label for ELEVIDYS; continued approval may be contingent upon verification of a clinical benefit in confirmatory trials; we may not be able to comply with all FDA requests in a timely manner or at all; the possible impact of regulations and regulatory decisions by the FDA and other regulatory agencies on our business, as well as the development of our product candidates and our financial and contractual obligations; our dependence on certain manufacturers to produce our products and product candidates, including any inability on our part to accurately anticipate product demand and to secure in a timely manner manufacturing capacity to meet product demand, may impair the availability of product to successfully support various programs; our data may not be sufficient for obtaining regulatory approval; if the actual number of patients living with Duchenne and LGMD is smaller than estimated, our revenue and ability to achieve profitability may be adversely affected; we are subject to uncertainty related to reimbursement policies; success in preclinical and clinical trials, especially if based on a small patient sample, does not ensure that later clinical trials will be successful, and the results of future research may not be consistent with past positive results or with advisory committee recommendations, or may fail to meet regulatory approval requirements for the safety and efficacy of product candidates; the commencement and completion of our clinical trials and announcement of results may be delayed or prevented for a number of reasons, including, among others, denial by the regulatory agencies of permission to proceed with our clinical trials, or placement of a clinical trial on hold, challenges in identifying, recruiting, enrolling and retaining patients to participate in clinical trials and inadeguate quantity or quality of supplies of a product candidate or other materials necessary to conduct clinical trials; different methodologies, assumptions and applications we use to assess particular safety or efficacy parameters may yield different statistical results, and even if we believe the data collected from clinical trials of our product candidates are positive, these data may not be sufficient to support approval by the FDA or other global regulatory authorities; we may not be able to execute on our business plans, including meeting our expected or planned regulatory milestones and timelines, research and clinical development plans, and bringing our product candidates to market, for various reasons, many of which may be outside of our control, including possible limitations of company financial and other resources, manufacturing limitations that may not be anticipated or resolved for in a timely manner, regulatory, court or agency decisions, such as decisions by the United States Patent and Trademark Office with respect to patents that cover our product candidates; and those risks identified under the heading "Risk Factors" in our most recent Annual Report on Form 10-K for the year ended December 31, 2022, and Form 10-Q filed with the Securities and Exchange Commission (SEC) as well as other SEC filings made by the Company, which you are encouraged to review

Any of the foregoing risks could materially and adversely affect the Company's business, results of operations and the trading price of Sarepta's common stock. For a detailed description of risks and uncertainties Sarepta faces, you are encouraged to review the SEC filings made by Sarepta. We caution investors not to place considerable reliance on the forward-looking statements contained in this presentation. Sarepta does not undertake any obligation to publicly update its forward-looking statements based on events or circumstances after the date hereof, except as required by law.

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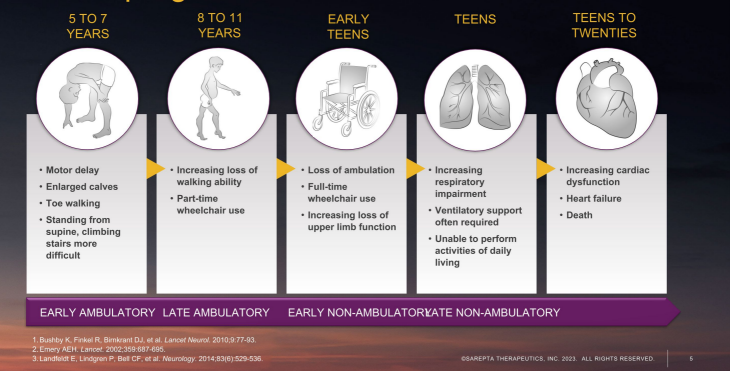
## **Elevidys** delandistrogene moxeparvovec-rokl

suspension for intravenous infusion

### An Historic Milestone In Genetic Medicine



## Disease progression in Duchenne<sup>1-3</sup>

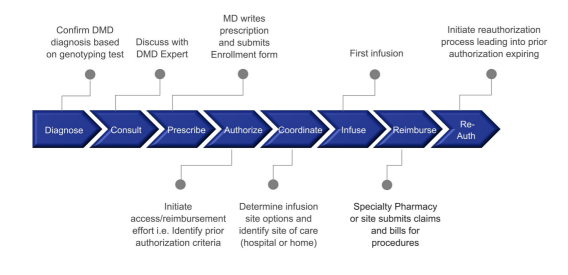


Sarepta is the recognized leader in Duchenne research and commercialization....





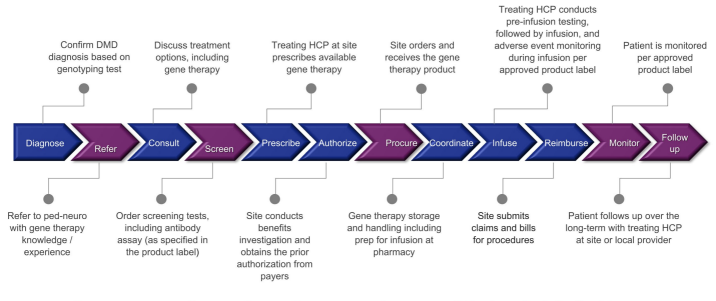
## PMO patient journey



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8

### Gene therapy patient journey...



### ... is more complex and requires engaging a multitude of stakeholders

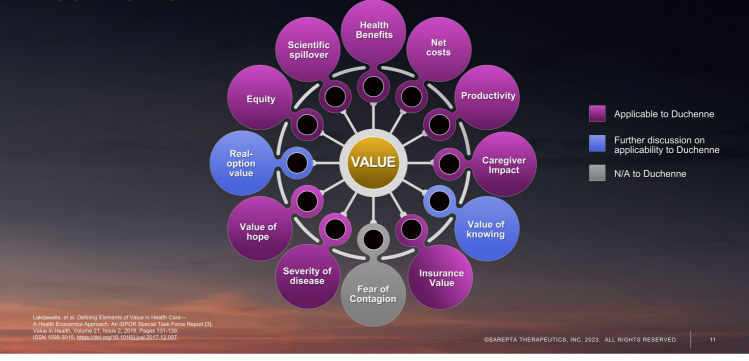
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## Journal of Market Access and Health Policy

Assessing the value of delandistrogene moxeparvovec (SRP-9001) gene therapy in patients with Duchenne muscular dystrophy in the United States

# A holistic approach to assessing innovative treatments

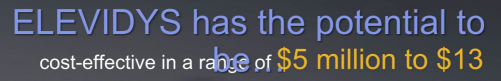


The cost-effectiveness analysis shows the value of ELEVIDYS in treating Duchenne

# ELEVIDYS has the potential to cost-effective in a rable of \$5 million to \$13

million compared to standard-of-care alone

The cost-effectiveness analysis shows the value of ELEVIDYS in treating Duchenne



million compared to standard-of-care alone

At a price of \$3.2 million ELEVIDYS is

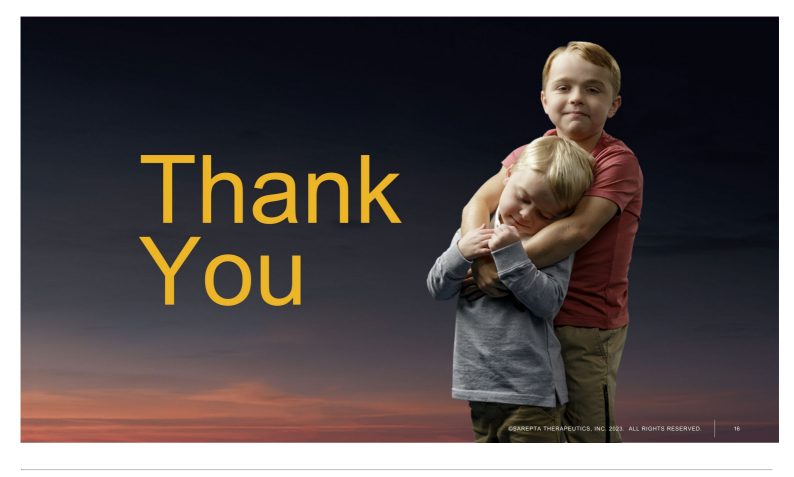
below the cost-effectiveness range

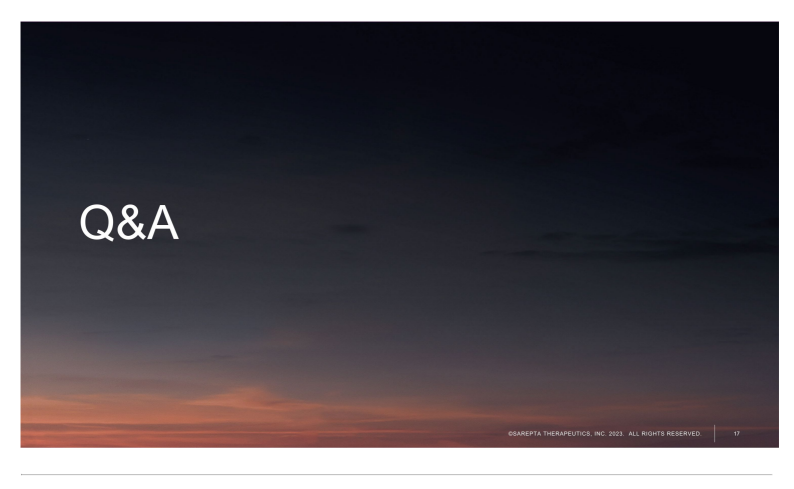
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Patients can't wait for the next breakthrough in medical research.

## So neither will we.

Doug Ingram President and CEO Sarepta Therapeutics, Inc. (NASDAQ: SRPT)

June 22, 2023



BENJAMIN Living with Duchenne muscular dystrophy