UNITED STATES SECURITIES AND EXCHANGE COMMISSION Washington, DC 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): September 13, 2021

Aquestive Therapeutics, Inc. (Exact name of Registrant as specified in its charter)

Delaware

(State or Other Jurisdiction of Incorporation or Organization)

001-38599 (Commission File Number)

82-3827296 (I.R.S. Employer Identification No.)

30 Technology Drive Warren, NJ 07059 (908) 941-1900

(Address, Including Zip Code, and Telephone Number, Including Area Code, of Registrant's Principal Executive Offices)

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 | AQST | Nasdaq Global 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 7.01 Regulation FD Disclosure.

Aquestive Therapeutics, Inc. (the "Company") is furnishing this Current Report on Form 8-K in connection with the disclosure of information, in the form of an investor presentation, to be given at meetings with institutional investors, analysts and others. This information may be amended or updated at any time and from time to time through another Current Report on Form 8-K, a later company filing or other means. A copy of the Company's investor presentation is attached hereto as Exhibit 99.1 to this Current Report on Form 8-K and incorporated into this Item 7.01 by reference and replaces in its entirety all prior investor presentations filed by the Company. The investor presentation is available on the Company's website located at www.aquestive.com, although the Company reserves the right to discontinue that availability at any time.

The information in this Item 7.01 (including Exhibit 99.1) shall not be deemed to be "filed" for purposes of, or otherwise subject to the liabilities of, Section 18 of the Securities Exchange Act of 1934, as amended (the "Exchange Act"), nor shall it be deemed to be incorporated by reference in any filing under the Securities Act of 1933, as amended, or the Exchange Act, except as shall be expressly set forth by specific reference in any such filing.

Item 9.01 Financial Statements and Exhibits

(d) Exhibits.

 Exhibit Number
 Description

 99.1
 Investor presentation dated September 2021.

SIGNATURE

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

Dated: September 13, 2021

Aquestive Therapeutics, Inc.

By: /s/ A. Ernest Toth, Jr

Name: A. Ernest Toth, Jr. Title: Chief Financial Officer



Aquestive Therapeutics Corporate Presentation

Advancing medicines. Solving problems. Improving lives.

Forward-Looking Statement

This presentation includes forward-looking statements within the meaning of the Private Securities Liligation Reform Act of 1995. Words such as "believe," "anticipate, "plan," "expect," "estimate," "intend," "may," "will," or the negative of those terms, and similar expressions, are intended to identify forward-looking statements. These forward-looking statements include, but are not limited to, statements megading the advancement of Libervant". AcQST-106, AQST-106, AQST-

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ation shall not constitute an offer to sell or the solicitation of an offer to buy these securities, nor shall there be any sale of these securities in any state or jurisdiction in which such offer, solicitation or sale would prior to registration or qualification under the securities laws of any such state or jurisdiction. wful prior to



COur Path

3

Proven track record Near-term pipeline Multiple cash-generating catalysts opportunities of success 10+ quarters of continuous Technology-based Libervant[™] Buccal Film (Diazepam) pharmaceutical company growth in commercial sales NDA filed on June 23, 2021 5 FDA-approved products PDUFA date December 23, 2021 Cash flow positive Epinephrine prodrug platform manufacturing business 10+ years of product sales with 2 clinical stage sublingual Business performance and capital 200+ patents worldwide • film candidates (SF) AQST-108 & options support commercial AQST-109 operations, Libervant launch and pipeline development

Libervant[®] Buccal Film (Diazepam) is an investigational drug being evaluated for use in children and adults with refractory seizures, who remain on stable regimens of antiepileptic drugs, to control bouts of increased seizure activity. The product profile, data from our trials, and related statements have not been approved by the FDA. Aquestive has received conditional acceptance of the use of this trade name, which is subject to final FDA review and acceptance.

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C PharmFilm[®] Technology – Where You Need It, When You Need It[™]





COur Team



Keith J. Kendall Chief Executive Officer and Director



Ken Marshall Chief Commercial Officer



Daniel Barber Chief Operating Officer



Mark Schobel Chief Innovation & Technology Officer



Peter Boyd SVP, Business Process & Information Technology



Gary H. Slatko, MD Chief Medical Officer, Neurology



Lori J. Braender General Counsel and Chief Compliance Officer



Ernie Toth Chief Financial Officer



Mark Lepore, MD Chief Medical Officer, Allergy



Theresa Wood SVP, Human Resources



C Our Products



C Solving problems in EPILEPSY:

SYMPAZAN Oral Film (Clobazam) LIBERVANT Buccal Film (Diazepam) Commercialized for Lennox-Gastaut syndrome 0 Under FDA review for management of refractory (LGS), a rare, severe form of epilepsy characterized by multiple manifestations of cognitive impairment and developmental delays¹. patients with epilepsy on stable regimens of AEDs who experience seizure clusters. ≈1.2M ≈700K Epilepsy patients⁴ suffer from uncontrolled, prescriptions annually refractory seizures ≈90% of patients with refractory seizures will not interact with the historically available treatment⁵ >1M Emergency Department 50K Of nearly 3.4MM 30-40% patients with epilepsy have seizures related to LGS² visits caused by seizures annually Of people with LGS have dysphagia³ C Aquestive 7 Advancing medici Solving problems. Improving lives.

C Continued Expansion of Epilepsy Franchise

SYMPAZAN: Sustained Script Growth for Over 10 Quarters

- Growing prescriber base, with over 30% penetration into the focused group of prescribers, with ~80% writing multiple scripts
- Launched as a precursor and compliment to Libervant
- 90% of focused group of prescribers write scripts for Sympazan and Libervant indications
- Solid two product sales force efficiency







C Solving Problems in ANAPHYLAXIS

- Anaphylaxis is an unpredictable, severe systemic allergic reaction that is rapid in onset and potentially fatal^7
- At-risk patients should always have immediate access to 2 doses of epinephrine⁸
- Delayed administration of epinephrine tied to increased fatalities⁹
- Aquestive is developing oral sublingual film formulations of epinephrine for treatment of allergic reactions (type 1), including anaphylaxis



C Key Pipeline Milestones



| 11 | Aquestive Advancing modelines. Solving problems. Improving lives. |
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C Financial Summary

Full Year 2021 Guidance (as of August 3, 2021)

- Total revenues of approximately \$46 million to \$48 million
- Non-GAAP adjusted gross margins of approximately 70% to 75% on total revenues
- Non-GAAP adjusted EBITDA loss of approximately \$39 million to \$42 million

Capital Adequacy

- Cash on hand (\$34.2 million at 6/30/21)
- Results from business performance, ATM activity, and expense management provide 12 months or more of capital with additional options and supports possible launch of Libervant and pipeline activities
- Debt reduced to \$51.5 million; potential additional capital, at Aquestive's option, of \$10 million after Libervant FDA approval and \$20 million after Libervant U.S. market access
- Available shelf registration



COur Focus in 2021

Continue to expand in our epilepsy franchise

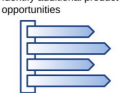
- Focus on the FDA approval and launch of LIBERVANT
- Generate continued growth in SYMPAZAN prescriptions



13

Advance our novel epinephrine delivery platform

- Complete and readout on clinical studies
- Request pre-IND meeting with FDA to establish path forward
- Identify additional product



Continue to strengthen the capital position of the Company

- Continued strong business performance to generate cash
- Access available funds on potential Libervant approval and U.S. market access
- Appropriate use of ATM facility
- Utilize shelf registration under favorable conditions



Advancing medicines Solving problems. Improving lives.



CORPORATE INFORMATION , PHARMFILM® TECHNOLOGY, SYMPAZAN®, LIBERVANT™ AND EPINEPHRINE DATA

1. van Rijckevorsel. Treatment of Lennox-Gastaut syndrome: overview and recent findings Neuropsychiatr Dis Treat. 2008;4(6):1001-1019 2. Kwan and Brodie. Early Identification of Refractory Epilepsy N Engl J Med. 2000;342(5):314-319

3. Ogawa K, Kanemoto K, Ishii Y, Koyama M, Sirasaka Y, Kawasaki J, Yamasaki S. Long-term follow-up study of Lennox–Gastaut syndrome in patients with severe motor and intellectual disabilities: with special reference to the problem of dysphagia. Seizure. 2001; 10:197-202

4. Laxer KD, Trinka E, Hirsch LJ, et al. The consequences of refractory epilepsy and its treatment. Epilepsy Behav. 2014;37:59-70.

5. Triangle Insights Group. Synthesis of Epilepsy (ARS) Primary Research. 2017. Internal Aquestive report: unpublished.

6. Pallin DJ, Goldstein JN, Moussally JS, Pelletier AJ, Green AR, Camargo CA Jr. Seizure visits in US emergency departments: epidemiology and potential disparities in care. Int J Emerg Med. 2008;1(2):97-105.

7. Simons F.E., Clark S., Camargo C.A. Jr: Anaphylaxis in the community: learning from the survivors. J Allergy Clin Immunol 2009, 124 (2): 301-306

8. Boyce JA, Assa'ad A, Burks AW, et al. Guidelines for the diagnosis and management of food allergy in the United States: report of the NIAID-sponsored expert panel. J Allergy Clin Immunol. 2010;126(6 Suppl):S1- S58.

9. Song TT, Lieberman P. Epinephrine in anaphylaxis: doubt no more. Current opinion in allergy and clinical immunology. 2015;15(4):323-8. 10. Shaker et al. (2020). Anaphylaxis—a 2020 practice parameter update, systematic review, and Grading of Recommendations, Assessment, Development and Evaluation (GRADE) analysis. J Allergy & Clin Immunology, Vol 143, Num 4 11. Yu J., Lin,R. The Epidemiology of Anaphylaxis. Clin Rev Allergy Immunol. 2018 Jun;54(3):366-374.doi: 10.1007/s12016-015-8503-x.

12. Borish, L., Danoff, T., Ma, L. VOLUME 133, ISSUE 2, SUPPLEMENT . doi.org/10.1016/j.jaci.2013.12.834

13. Symphony Health 2020 data on file.



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Thank You

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