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): March 07, 2022

KALVISTA PHARMACEUTICALS, INC.

(Exact name of Registrant as Specified in Its Charter)

Delaware (State or Other Jurisdiction of Incorporation) 001-36830 (Commission File Number) 20-0915291 (IRS Employer Identification No.)

55 Cambridge Parkway Suite 901E Cambridge, Massachusetts (Address of Principal Executive Offices)

02142 (Zip Code)

Registrant's Telephone Number, Including Area Code: 857 999-0075

(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)) П 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, \$0.001 par value per share KALV The NASDAQ Stock Market LLC 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 \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. \square

Item 7.01 Regulation FD Disclosure.

On March 7, 2022, the Company issued a press release announcing that it has initiated the Phase 3 KONFIDENT clinical trial evaluating the safety and efficacy of KVD900 as the first potential oral, on-demand therapy for hereditary angioedema (HAE) attacks. A copy of the press release is furnished with this Current Report on Form 8-K as Exhibit 99.1.

Item 8.01 Other Events.

On March 7, 2022, the Company announced that it has initiated the Phase 3 KONFIDENT clinical trial evaluating the safety and efficacy of KVD900 as the first potential oral, on-demand therapy for hereditary angioedema (HAE) attacks.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits.

Exhibit

| <u>Number</u> | <u>Description</u> |
|---------------|--|
| 99.1 | Press release dated March 7, 2022 |
| 104 | Cover Page Interactive Data File (embedded within the Inline XBRL document). |

Forward-Looking Statements

This Current Report on Form 8-K contains "forward-looking" statements within the meaning of the safe harbor provisions of the U.S. Private Securities Litigation Reform Act of 1995. Forward-looking statements can be identified by words such as: "anticipate," "intend," "plan," "goal," "seek," "believe," "project," "estimate," "expect," "strategy," "future," "likely," "may," "should," "will" and similar references to future periods. These statements are subject to numerous risks and uncertainties, including the potential impact of the ongoing COVID-19 pandemic, that could cause actual results to differ materially from what we expect. Examples of forward-looking statements include, among others, timing or outcomes of communications with the FDA, our expectations about safety and efficacy of our product candidates and timing of clinical trials and its results, our ability to commence clinical studies or complete ongoing clinical studies, including our Phase 3 KONFIDENT and Phase 2 KOMPLETE clinical trials, and to obtain regulatory approvals for KVD900, KVD824 and other candidates in development, the ability of KVD900, KVD824 and other candidates in development to treat HAE or DME, and the future progress and potential success of our oral Factor XIIa program. Further information on potential risk factors that could affect our business and financial results are detailed in our filings with the Securities and Exchange Commission, including in our annual report on Form 10-K for the year ended April 30, 2021, our quarterly reports on Form 10-Q, and our other reports that we may make from time to time with the Securities and Exchange Commission. We undertake no obligation to publicly update any forward-looking statement, whether written or oral, that may be made from time to time, whether as a result of new information, future developments or otherwise

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.

KALVISTA PHARMACEUTICALS, INC.

Date: March 7, 2022

/s/ Benjamin L. Palleiko Benjamin L. Palleiko Chief Business Officer and Chief Financial Officer

KalVista Pharmaceuticals Announces Initiation of KVD900 Phase 3 KONFIDENT Clinical Trial

-KONFIDENT to Evaluate KVD900 As First Oral On-Demand Therapy for HAE-

-KONFIDENT Designed to Support Broad Label for Treatment of All HAE Attacks-

Cambridge, MA and Salisbury, England, March 7, 2022 – KalVista Pharmaceuticals, Inc. (NASDAQ: KALV), a clinical stage pharmaceutical company focused on the discovery, development, and commercialization of oral, small molecule protease inhibitors, today announced the initiation of the Phase 3 KONFIDENT clinical trial evaluating the efficacy and safety of KVD900 as the first potential oral, on-demand therapy for hereditary angioedema (HAE) attacks. This worldwide, double-blind, placebo-controlled crossover trial will evaluate the efficacy of two dose levels of KVD900 compared to placebo in adolescents and adults experiencing acute HAE attacks. KVD900 is the most advanced potential oral on-demand therapy for HAE in clinical development, and is intended to provide a substantial improvement over the current on-demand therapies for HAE attacks, which are all delivered by injection.

"Beginning the KONFIDENT trial represents a major milestone for KalVista," said Andrew Crockett, Chief Executive Officer of KalVista. "We believe that KVD900 has the potential to transform the treatment paradigm for HAE patients experiencing acute attacks, whether they primarily treat with on-demand medications or use long-term prophylaxis. Based upon the results of our Phase 2 study released last year, we expect that KVD900 can provide patients with symptom relief as rapidly as existing therapies, but with an oral tablet that will allow earlier treatment of all patient-recognized HAE attacks. Our goal is to provide patients with the confidence that their attacks will be controlled in the earliest stages and without the associated treatment pain and other challenges of injectable therapies."

The Phase 3 KONFIDENT trial is a worldwide clinical study being conducted at approximately 60 sites in 20 countries. The trial is intended to enroll a minimum of 84 HAE adolescent and adult patients who will complete treatment of three attacks: one each with 300 mg KVD900, 600 mg KVD900 and placebo in a double-blinded, randomized sequence. The primary endpoint of the trial is time to the beginning of symptom relief, evaluated on a Patient Global Impression of Change (PGI-C) scale, and additional endpoints will evaluate other measures of patient response and attack progression, as well as safety. Patients will dose upon first recognition of an attack, and all attack types including laryngeal attacks will be eligible for treatment. Patients will be permitted to take an additional dose of investigational drug, if symptoms warrant, and will always have access to their conventional injectable therapy. Study participants also will be allowed to maintain their prophylaxis regimen if they were receiving one at study enrollment.

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KalVista currently anticipates that data from KONFIDENT will be available in the second half of 2023 and will provide further updates as the trial progresses. Additional information about KONFIDENT can be found at www.konfidentstudy.com and www.clinicaltrials.gov.

About KalVista Pharmaceuticals, Inc.

KalVista Pharmaceuticals, Inc. is a pharmaceutical company focused on the discovery, development, and commercialization of small molecule protease inhibitors for diseases with significant unmet need. KalVista has developed a proprietary portfolio of novel, small molecule plasma kallikrein inhibitors initially targeting hereditary angioedema (HAE) and diabetic macular edema (DME). KalVista is developing KVD900 as an oral on-demand therapy for acute HAE attacks and has initiated the Phase 3 KONFIDENT clinical trial. KVD824 is in development for prophylactic treatment of HAE, with the Phase 2 KOMPLETE clinical trial underway. In addition, KalVista's oral Factor XIIa inhibitor program represents a new generation of therapies that may further improve the treatment of HAE for patients. In DME, an intravitreally administered plasma kallikrein inhibitor, called KVD001, has completed a Phase 2 clinical trial.

For more information about KalVista, please visit www.kalvista.com.

For more information on the KVD900 HAE on-demand Phase 3 KONFIDENT study, please visit www.konfidentstudy.com_ For more information on the KVD824 HAE prophylaxis Phase 2 KOMPLETE study, please visit www.kompletestudy.com_

Forward-Looking Statements

This press release contains "forward-looking" statements within the meaning of the safe harbor provisions of the U.S. Private Securities Litigation Reform Act of 1995. Forward-looking statements can be identified by words such as: "anticipate," "intend," "plan," "goal," "seek," "believe," "project," "estimate," "expect," "strategy," "future," "likely," "may," "should," "will" and similar references to future periods. These statements are subject to numerous risks and uncertainties, including the potential impact of the ongoing COVID-19 pandemic, that could cause actual results to differ materially from what we expect. Examples of forward-looking statements include, among others, timing or outcomes of communications with the FDA, our expectations about safety and efficacy of our product candidates and timing of clinical trials and its results, our ability to commence clinical studies or complete ongoing clinical studies, including our Phase 3 KONFIDENT and Phase 2 KOMPLETE clinical trials, and to obtain regulatory approvals for KVD900, KVD824 and other candidates in development, the ability of KVD900, KVD824 and other candidates in development to treat HAE or DME, and the future progress and potential success of our oral Factor XIIa program. Further information on potential risk factors that could affect our business and financial results are detailed in our filings with the Securities and Exchange Commission, including in our annual report on Form 10-K for the year ended April 30, 2021, our quarterly reports on Form 10-Q, and our other reports that we may

make from time to time with the Securities and Exchange Commission. We undertake no obligation to publicly update any forward-looking statement, whether written or oral, that may be made from time to time, whether as a result of new information, future developments or otherwise.

Contact:

KalVista Pharmaceuticals, Inc. Ben Palleiko CBO & CFO 857-999-0890 investors@kalvista.com