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 13, 2024



NEUROBO PHARMACEUTICALS, INC.

(Exact name of Registrant as Specified in Its Charter)

Delaware	001-37809	47-2389984	
(State or other jurisdiction	(Commission	(IRS Employer	
of incorporation)	File Number)	Identification No.)	
545 Concord Avenue, Suite 210 Cambridge, Massachusetts			
		02138	
(Address of principal executive offices)		(Zip Code)	
	(857) 702-9600		
(Registrant's to	elephone number, inclu	iding area code)	
	Not applicable		
(Former name or fo	ormer address, if chang	ged since last report)	
Check the appropriate box below if the Form 8-K registrant under any of the following provisions:	filing is intended to si	multaneously satisfy the filing obligation of the	
 □ 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:		
	Trading		
Title of each class	Symbol(s)	Name of each exchange on which registered	
Common Stock, par value \$0.001 per share	NRBO	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 □			
If an emerging growth company, indicate by check period for complying with any new or revised fine Exchange Act. □			

Item 7.01. Regulation FD Disclosure.

On August 13, 2024, NeuroBo Pharmaceuticals, Inc. (the "Company") issued a press release announcing the completion of enrollment of the single ascending dose (SAD) Part 1 of its Phase 1 clinical trial of DA-1726, a novel, dual oxyntomodulin (OXM) analog agonist that functions as a glucagon-like peptide-1 receptor (GLP1R) and glucagon receptor (GCGR), for the treatment of obesity. A total of 45 participants have been enrolled and randomized into one of 5 cohorts, with each cohort having been randomized in a 6:3 ratio of DA-1726 or placebo. A copy of the press release is furnished as Exhibit 99.1 to this Current Report on Form 8-K (the "Report") and incorporated herein by reference.

Information contained on or accessible through any website reference in the press release is not part of, or incorporated by reference in, this Report, and the inclusion of such website addresses in this Report by incorporation by reference of the press release is as inactive textual references only.

Exhibit 99.1 hereto contains forward-looking statements within the meaning of the federal securities laws. These forward-looking statements are based on current expectations and are not guarantees of future performance. Further, the forward-looking statements are subject to the limitations listed in Exhibit 99.1 and in the other reports of the Company filed with the Securities and Exchange Commission, including that actual events or results may differ materially from those in the forward-looking statements.

The information in this Report, including Exhibit 99.1 hereto, is furnished pursuant to Item 7.01 and 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 liabilities 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. The Company's submission of this Report shall not be deemed an admission as to the materiality of any information required to be disclosed solely to satisfy the requirements of Regulation FD.

Item 9.01. Financial Statements and Exhibits.

(d) Exhibits

Exhibit	
Number	Exhibit Description
99.1	Press Release dated August 13, 2024.
104	Cover Page Interactive Data File (embedded within 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.

NEUROBO PHARMACEUTICALS, INC.

Date: August 13, 2024 By: /s/ Hyung Heon Kim

Hyung Heon Kim

President and Chief Executive Officer



NeuroBo Pharmaceuticals Completes Enrollment of the SAD Part 1 of Its Phase 1 Clinical Trial Evaluating DA-1726 for the Treatment of Obesity

No Significant Issues Encountered During the Single Ascending Dose (SAD) Study, Allowing for Previously
Reported Accelerated Start to Multiple Ascending Dose (MAD) Part 2

Top Line Data Readout from SAD Part 1 Expected in the Third Quarter of 2024, and from the MAD Part 2 in the First Quarter of 2025

CAMBRIDGE, Mass., August 13, 2024 – NeuroBo Pharmaceuticals, Inc. (Nasdaq: NRBO), a clinical-stage biotechnology company focused on transforming cardiometabolic diseases, today announced the completion of enrollment of the single ascending dose (SAD) Part 1 of its Phase 1 clinical trial of DA-1726, a novel, dual oxyntomodulin (OXM) analog agonist that functions as a glucagon-like peptide-1 receptor (GLP1R) and glucagon receptor (GCGR), for the treatment of obesity. A total of 45 participants have been enrolled and randomized into one of 5 cohorts, with each cohort having been randomized in a 6:3 ratio of DA-1726 to placebo.

"Completion of enrollment in Part 1 of this Phase 1 clinical trial evaluating DA-1726 for the treatment of obesity marks the achievement of yet another key milestone for NeuroBo and reflects our ongoing commitment to transforming the treatment of cardiometabolic diseases with differentiated therapies," stated Hyung Heon Kim, President and Chief Executive Officer of NeuroBo. "Notably, we encountered no significant issues during the SAD study, allowing us to begin the multiple ascending dose (MAD) study ahead of schedule. As previously reported, strong pre-clinical data has shown superior weight loss with DA-1726 versus semaglutide (Wegovy®) and similar weight reduction while consuming more food compared to tirzepatide (Zepbound®). Data presented at the American Diabetes Association 84th Scientific Sessions showed that DA-1726 demonstrated superior weight loss compared to survodutide, a drug with the same mechanism of action, while also demonstrating retention of relative lean body mass preservation compared to survodutide while also exhibiting superior glucose lowering. These factors lead us to believe that DA-1726 may eventually become a best-in-class obesity drug with a better tolerability profile than currently marketed GLP-1 agonists, and those now in late-stage clinical trials, based on its balanced activation of GLP1R and glucagon receptors, while increasing energy expenditure."

Mr. Kim continued, "Our estimated timelines for this program remain unchanged. We expect to report top-line data from the SAD Part 1 portion of the Phase 1 clinical trial in the third quarter of this year and top-line data from the MAD Part 2 in the first quarter of 2025. Further, upon clearance of an updated Investigational New Drug (IND) application with the U.S. Food and Drug Administration (FDA), we expect to dose the first patient in the planned Part 3 of the trial during the third quarter of 2025, providing an interim data readout in or around mid-2026 and issuing top-line results in the second half of 2026," concluded Mr. Kim.

The Phase 1 trial is currently designed to be a randomized, placebo-controlled, double-blind study to investigate the safety, tolerability, pharmacokinetics (PK), and pharmacodynamics (PD) of single and multiple ascending doses of DA-1726 in obese, otherwise healthy subjects. Part 2, currently enrolling

subjects, is designed as a MAD study, and is expected to enroll approximately 36 participants, who will be randomized at the same 6:3 ratio into 4 planned cohorts, each to receive 4 weekly administrations of DA-1726 or placebo. The first patient in the MAD study was dosed ahead of schedule, in late June, as previously reported.

The primary endpoint of the Phase 1 trial will assess the safety and tolerability of DA-1726 by monitoring adverse events (AEs), serious adverse events (SAEs), treatment emergent adverse events (TEAEs) and AEs leading to treatment discontinuation. Secondary endpoints include the PK of DA-1726, assessed via serum concentrations over time and metabolite profiling at the highest doses of DA-1726. Exploratory endpoints will include the effect of DA-1726 on metabolic parameters, cardiac parameters, fasting lipid levels, body weight, waist circumference and body mass index (BMI), among others.

For more information on this clinical trial, please visit: www.clinicaltrials.gov NCT06252220.

About DA-1726

DA-1726 is a novel oxyntomodulin (OXM) analogue functioning as a GLP1R/GCGR dual agonist for the treatment of obesity and Metabolic Dysfunction-Associated Steatohepatitis (MASH) that is to be administered once weekly subcutaneously. DA-1726 acts as a dual agonist of GLP-1 receptors (GLP1R) and glucagon receptors (GCGR), leading to weight loss through reduced appetite and increased energy expenditure. DA-1726 has a well understood mechanism and, in pre-clinical mice models, resulted in improved weight loss compared to semaglutide and cotadutide (another OXM analogue). Additionally, in pre-clinical mouse models, DA-1726 elicited similar weight reduction, while consuming more food, compared tirzepatide and survodutide, while also preserving lean body mass and demonstrating improved lipid-lowering effects compared to survodutide.

About NeuroBo Pharmaceuticals

NeuroBo Pharmaceuticals, Inc. is a clinical-stage biotechnology company focused on transforming cardiometabolic diseases. The company is currently developing DA-1726 for the treatment of obesity, and is developing DA-1241 for the treatment of Metabolic Dysfunction-Associated Steatohepatitis (MASH). DA-1726 is a novel oxyntomodulin (OXM) analogue that functions as a glucagon-like peptide-1 receptor (GLP1R) and glucagon receptor (GCGR) dual agonist. OXM is a naturally-occurring gut hormone that activates GLP1R and GCGR, thereby decreasing food intake while increasing energy expenditure, thus potentially resulting in superior body weight loss compared to selective GLP1R agonists. DA-1241 is a novel G-protein-coupled receptor 119 (GPR119) agonist that promotes the release of key gut peptides GLP-1, GIP, and PYY. In pre-clinical studies, DA-1241 demonstrated a positive effect on liver inflammation, lipid metabolism, weight loss, and glucose metabolism, reducing hepatic steatosis, hepatic inflammation, and liver fibrosis, while also improving glucose control.

For more information, please visit www.neurobopharma.com.

Forward Looking Statements

Certain statements in this press release may be considered forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995. Words such as "believes", "expects", "anticipates", "may", "will", "should", "seeks", "approximately", "potential", "intends", "projects", "plans", "estimates" or the negative of these words or other comparable terminology (as well as other words or expressions referencing future events, conditions or circumstances) are intended to identify forward-looking statements. Forward-looking statements are predictions, projections and other

statements about future events that are based on current expectations and assumptions and, as a result, are subject to risks and uncertainties. Many factors could cause actual future events to differ materially from the forward-looking statements in this press release, including, without limitation, those risks associated with NeuroBo's ability to execute on its commercial strategy; the timeline for regulatory submissions; the ability to obtain regulatory approval through the development steps of NeuroBo's current and future product candidates; the ability to realize the benefits of the license agreement with Dong-A ST Co. Ltd., including the impact on future financial and operating results of NeuroBo; the cooperation of NeuroBo's contract manufacturers, clinical study partners and others involved in the development of NeuroBo's current and future product candidates; potential negative interactions between NeuroBo's product candidates and any other products with which they are combined for treatment; NeuroBo's ability to initiate and complete clinical trials on a timely basis; NeuroBo's ability to recruit subjects for its clinical trials; whether NeuroBo receives results from NeuroBo's clinical trials that are consistent with the results of pre-clinical and previous clinical trials; impact of costs related to the license agreement, known and unknown, including costs of any litigation or regulatory actions relating to the license agreement; the effects of changes in applicable laws or regulations; the effects of changes to NeuroBo's stock price on the terms of the license agreement and any future fundraising; and other risks and uncertainties described in NeuroBo's filings with the Securities and Exchange Commission, including NeuroBo's most recent Annual Report on Form 10-K. Forward-looking statements speak only as of the date when made. NeuroBo does not assume any obligation to publicly update or revise any forwardlooking statements, whether as a result of new information, future events or otherwise, except as required by law.

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