UNITED STATES SECURITIES AND EXCHANGE COMMISSION Washington, D.C. 20549

	FORM 8-K	
	CURRENT REPORT	
Pursuant to Sect	ion 13 or 15(d) of the Securities Exc	change Act of 1934
Date of Rep	oort (Date of earliest event reported): Nov	ember 2, 2023
		_
(E	ALIGOS THERAPEUTICS, INC	
Delaware (State or Other Jurisdiction of Incorporation)	001-39617 (Commission File Number)	82-4724808 (I.R.S. Employer Identification No.)
(A	One Corporate Dr., 2nd Floor South San Francisco, California 94080 ddress of Principal Executive Offices) (Zip C	Code)
(R	(800) 466-6059 egistrant's telephone number, including area	code)
(Forme	er name or former address, if changed since la	ist report)
Check the appropriate box below if the Form 8-K filin following provisions:	g is intended to simultaneously satisfy the fil	— ing obligation of the registrant under any of the
 □ Written communications pursuant to Rule 425 und □ Soliciting material pursuant to Rule 14a-12 under □ Pre-commencement communications pursuant to □ Pre-commencement communications pursuant to 	the Exchange Act (17 CFR 240.14a-12) Rule 14d-2(b) under the Exchange Act (17 C	
Securities registered pursuant to Section 12(b) of the $\it A$	Act:	
Title of each class	Trading Symbol(s) ALGS	Name of each exchange on which registered
Common Stock, \$0.0001 par value per share Indicate by check mark whether the registrant is an emchapter) or Rule 12b-2 of the Securities Exchange Act	nerging growth company as defined in Rule 4	The Nasdaq Stock Market LLC 05 of the Securities Act of 1933 (§230.405 of this
Emerging growth company ⊠		
If an emerging growth company, indicate by check ma or revised financial accounting standards provided pur		extended transition period for complying with any new □

Item 2.02. Results of Operations and Financial Condition.

On November 2, 2023, the Registrant issued a press release, a copy of which is attached hereto as Exhibit 99.1 and is incorporated herein by reference.

The information in this Item 2.02 and the attached Exhibit 99.1 are being furnished and shall not be deemed to be "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, nor shall they be deemed to be incorporated by reference in any filing made by the Registrant under the Securities Act of 1933, as amended, or the Exchange Act, except as shall be expressly set forth by specific reference in such a filing.

Item 9.01. Financial Statements and Exhibits.

99.1 Press Release dated November 2, 2023

104

Cover Page Interactive Data File (embedded within the Inline XBRL document)

SIGNATURE

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.

Aligos Therapeutics, Inc.

Date: November 2, 2023 By: /s/ Lesley Ann Calhoun

Lesley Ann Calhoun

Executive Vice President, Chief Financial Officer

Aligos Therapeutics Reports Recent Business Progress and Third Quarter 2023 Financial Results

SOUTH SAN FRANCISCO, Calif., Nov. 02, 2023 (GLOBE NEWSWIRE) -- Aligos Therapeutics, Inc. (Nasdaq: ALGS), a clinical stage biopharmaceutical company focused on developing novel therapeutics to address unmet medical needs in liver and viral diseases, today reported recent business progress and financial results for the third quarter 2023.

"Over this quarter, we raised approximately \$92 million in a private placement financing allowing the advancement of our NASH THR-ß and CHB CAM-E programs. Additionally, we successfully received clearance for Aligos' first US IND (for ALG-055009) and secured an \$8.5 million contract with the NIAID to advance our coronavirus protease inhibitor into Phase 2 clinical trials. We are proud to have achieved these critical milestones," noted Lawrence Blatt, Ph.D., MBA, Chairman & CEO of Aligos Therapeutics. "With these achievements in hand, our team is well positioned to execute on our priorities, which include conducting a Ph2a MRI-PDFF study in NASH with ALG-055009, conducting Phase 2 enabling activities for our promising CAM-E, ALG-000184, in CHB, and advancing our potentially best-in-class coronavirus protease inhibitor, ALG-097558. Over the next quarter our research team will continue to identify promising novel small molecule drug candidates as backup molecules for our lead clinical programs as well as discover new small molecules targeting novel molecular pathways in viral and liver diseases."

Recent Business Progress

Aligos Portfolio of Drug Candidates

NASH (THR-ß; ALG-055009)

- The US IND was cleared by the FDA, which now enables execution of a planned Phase 1 statin drug-drug interaction study in Q4 2023, prior to filing the Phase 2a protocol to the IND in Q4 2023
- In the meantime, Phase 2a startup activities were initiated and are now underway
- A preliminary Phase 2a study design was formulated in close collaboration with the study's Principal Investigator, Dr. Stephen Harrison. The final study design is subject to feedback from the FDA which is expected in early Q1 2024
- The final Phase 1 FIH study data were accepted as a poster presentation (Poster #2900-A) at the AASLD Liver Meeting in November 2023

CHB (CAM-E, ALG-000184)

- Dosing and enrollment in ongoing Study ALG-000184-201 continues. The total dosing duration now exceeds 48 weeks in some subjects in this Phase 1a/1b study
- Data from long term dosing cohorts in Study ALG-000184-201, including a late breaking poster (Poster #5028-C) summarizing the effects of ALG-000184 on various viral markers, will be presented at the AASLD Liver Meeting in November 2023
- With the recent completion of the PIPE, we plan to initiate Ph2 enabling activities, including manufacturing of drug supply, in the near future

Coronavirus (PI, ALG-097558)

- Dosing continues in the ongoing first in human study, ALG-097558-701. Dosing is ongoing in the single ascending dose portion of the study and is due to start soon in the multiple ascending dose portion. We continue to project having topline data from this important safety and pharmacokinetics study in H1 2024
- The NIAID awarded Aligos an \$8.5M contract to conduct Phase 2 enabling activities, which includes multiple nonclinical and clinical studies. These studies will be initiated in H1 2024 and topline data are expected in H2 2025. The ALG-097558 program has been awarded over \$11M across two NIH grants and contracts and we plan to seek additional external funding from both public and private sources to further advance this important program

Financial Results for the Third Quarter 2023

Cash, cash equivalents and investments totaled \$70.4 million as of September 30, 2023, compared with \$125.8 million as of December 31, 2022. Additionally, we raised in private placement financing approximately \$92 million in gross proceeds, before deducting placement agent's fees and other expenses, in October 2023. Including the expected net proceeds from the private placement, we believe our cash balance provides sufficient cash to fund planned operations through the end of 2025.

Net losses for the three months ended September 30, 2023 were \$18.0 million or basic and diluted net loss per common share of \$(0.41), compared to net losses of \$18.6 million or basic and diluted net loss per common share of \$(0.44) for the three months ended September 30, 2022.

Research and development (R&D) expenses for the three months ended September 30, 2023 were \$15.9 million, compared with \$17.8 million for the same period of 2022. The decrease was primarily due to employee-related costs and other costs including facility expenses, partially offset by an increase in third party expenses due to the milestone payments made as a result of dosing the first patient in a clinical trial. Total R&D stock-based compensation expense incurred for the three months ended September 30, 2023 was \$1.6 million, compared with \$1.9 million for the same period of 2022.

General and administrative (G&A) expenses for the three months ended September 30, 2023 were \$6.4 million, compared with \$5.3 million for the same period of 2022. The increase in G&A expenses for this comparative period is primarily attributable to an increase in legal and related costs. Total G&A stock-based compensation expense incurred for the three months ended September 30, 2023 was \$1.6 million, compared with \$1.5 million for the same period of 2022.

About Aligos

Aligos Therapeutics, Inc. is a clinical stage biopharmaceutical company that was founded in 2018 with the mission to become a world leader in the treatment of liver and viral diseases. Aligos' strategy is to harness the deep expertise and decades of drug development experience its team has in liver and viral diseases to discover and develop potentially best-in-class therapeutics for nonalcoholic steatohepatitis (NASH) and viruses with high unmet medical need such as coronaviruses and chronic hepatitis B (CHB).

Forward-Looking Statement

This press release contains forward-looking statements within the meaning of the U.S. Private Securities Litigation Reform Act of 1995. Any statements in this press release that are not historical facts may be considered "forward-looking statements," including without limitation, statements that the team is well positioned to execute on the company's priorities which include conducting a Ph2a MRI-PDFF study in NASH with ALG-055009, conducting Phase 2 enabling activities for the CAM-E, ALG-000184, in CHB, and advancing its coronavirus protease inhibitor, ALG-097558, and that over the next quarter, the research team will continue to identify promising novel small molecule drug candidates as backup molecules for the lead clinical programs as well as discover new small molecules targeting novel molecular pathways in viral and liver diseases; with respect to the NASH ALG-055009 program, that the company plans to execute on a Phase 1 statin drug-drug interaction study in Q4 2023 prior to filing the Phase 2a protocol to the IND in Q4 2023, with final study design subject to feedback from the FDA, expected in early Q1 2024; with respect to the CHB ALG-000184 program, that the dosing and enrollment in ongoing Study ALG-000184-201 continues and the plan is to initiate Ph2 enabling activities, including manufacturing of drug supply, in the near future; with respect to the company's COVID-19 ALG-097558 program, that dosing continues in the single ascending dose portion of the ALG-097558-701 study and is due to start soon in the multiple ascending dose portion, that the company continues to project having topline data from this safety and pharmacokinetics study in H1 2024, that the plan is to initiate Phase 2 enabling activities, which include multiple nonclinical and clinical studies, in H1 2024, that the expectation is to have topline data in H2 2025 and that the plan is to seek additional external funding from both public and private sources to further advance the program; and that the company continues to believe its cash balance provides sufficient cash to fund planned operations through the end of 2025. Forward-looking statements are typically, but not always, identified by the use of words such as "may," "will," "would," "believe," "intend," "plan," "anticipate," "estimate," "expect," and other similar terminology indicating future results. Such forward looking statements are subject to substantial risks and uncertainties that could cause our development programs, future results, performance, or achievements to differ materially from those anticipated in the forward-looking statements. Such risks and uncertainties include, without limitation, risks and uncertainties inherent in the drug development process, including Aligos' clinical-stage of development, the process of designing and conducting clinical trials, the regulatory approval processes, the timing of regulatory filings, the challenges associated with manufacturing drug products, Aligos' ability to successfully establish, protect and defend its intellectual property, other matters that could affect the sufficiency of Aligos' capital resources to fund operations, reliance on third parties for manufacturing and development efforts, and the impact of global events and other macroeconomic conditions on the Aligos' business. For a further description of the risks and uncertainties that could cause actual results to differ from those anticipated in these forward-looking statements, as well as risks relating to the business of Aligos in general, see Aligos' Quarterly Report on Form 10-Q filed with the Securities and Exchange Commission on November 2, 2023 and its future periodic reports to be filed or submitted with the Securities and Exchange Commission. Except as required by law, Aligos undertakes no obligation to update any forward-looking statements to reflect new information, events or circumstances, or to reflect the occurrence of unanticipated events.

Aligos Therapeutics, Inc Condensed Consolidated Statements of Operations (In thousands, except share and per share amounts)

		Three Months Ended September 30,				Nine Months Ended				
						Septe	er 30,			
		2023		2022		2023		2022		
		(Unaudited)	•'	(Unaudited)	- '-	(Unaudited)		(Unaudited)		
Revenue from Collaborations	\$	2,154	\$	4,106	\$	7,329	\$	10,370		
Revenue from Customers		1,085		-		5,519		-		
Operating Expenses:										
Research and development		15,867		17,791		50,783		65,977		
General and administrative		6,443		5,263		24,195		19,291		
Total operating expenses		22,310		23,054		74,978		85,268		
Loss from operations		(19,071)		(18,948)		(62,130)		(74,898)		

Interest and other income, net Loss before income tax expense	_	1,059 (18,012)	_	284 (18,664)	_	3,168 (58,962)	_	795 (74,103)
Income tax benefit (expense) Net loss Basic and diluted net loss per common share	_ = \$_	(29) (18,041) (0.41)	- - \$_	43 (18,621) (0.44)	\$	(825) (59,787) (1.38)	- - \$_	(57) (74,160) (1.74)
Weighted-average number of shares used in computing basic and diluted net loss per common share	_	43,496,975	=	42,761,928		43,209,656	_	42,647,732

Aligos Therapeutics, Inc. Condensed Consolidated Balance Sheets (In thousands)

	September 30, 2023			December 31, 2022		
	_	(Unaudited)	_	(audited) (1)		
Assets						
Current assets:						
Cash and cash equivalents	\$	70,429	\$	81,347		
Short-term investments		-		44,480		
Prepaid expenses and other current assets		4,214		7,718		
Total current assets		74,643		133,545		
Other assets		11,209		13,148		
Total assets	\$	85,852	\$	146,693		
Liabilities and Stockholders' Equity						
Current liabilities	\$	22,468	\$	33,129		
Other liabilities, noncurrent		8,563		9,664		
Total liabilities		31,031		42,793		
Total stockholders' equity		54,821		103,900		
Total liabilities and stockholders' equity	\$	85,852	\$	146,693		

⁽¹⁾ The balance sheet as of December 31, 2022 has been derived from the audited consolidated financial statements at that date included in the Company's Annual Report on Form 10-K for the year ended December 31, 2022.

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Investor Contact

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