## 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 9, 2021

### Allakos Inc.

(Exact name of registrant as specified in its charter)

Delaware (State or other jurisdiction of incorporation) 001-38582 (Commission File Number) 45-4798831 (IRS Employer Identification No.)

975 Island Drive, Suite 201
Redwood City, California 94065

(Address of principal executive offices, including zip code)

(650) 597-5002

 $(Registrant's\ telephone\ number,\ including\ area\ code)$ 

Not Applicable

(Former name or former address, if changed since last report)

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	ALLK	The Nasdaq Global Select Market					
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 (see General Instruction A.2. below):								
	□ 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))							
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. $\Box$								

#### Item 2.02 Results of Operations and Financial Condition.

On August 9, 2021, Allakos Inc. (the "Company") issued a press release reporting its financial results for the second quarter ended June 30, 2021. The full text of the press release is attached as Exhibit 99.1 to this Current Report on Form 8-K and is incorporated herein by reference.

The information in this Item 2.02 of this Form 8-K, including the attached Exhibit 99.1, is intended to be furnished 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, nor shall it be deemed incorporated by reference in any filing made by the Company under the Securities Act of 1933, as amended, or the Exchange Act, except as expressly set forth by specific reference in such filing.

#### Item 9.01 Financial Statements and Exhibits.

(d) Exhibits

Exhibit Number	Description
99.1	Press Release dated August 9, 2021.
104	Cover Page Interactive Data File (embedded within the Inline XBRL document)
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#### **SIGNATURES**

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.

Date: August 9, 2021

Allakos Inc.

By: /s/ Robert Alexander

Robert Alexander

Chief Executive Officer



#### Allakos Provides Business Update and Reports Second Quarter 2021 Financial Results

**REDWOOD CITY, Calif., August 9, 2021** – Allakos Inc. (the "Company") (Nasdaq: ALLK), a biotechnology company developing lirentelimab (AK002) for the treatment of eosinophil and mast cell-related diseases, today provided a business update and reported financial results for the second quarter ended June 30, 2021.

#### **Recent Accomplishments**

- Presented new data at Digestive Disease Week (DDW) 2021 suggesting eosinophilic gastritis and/or eosinophilic duodenitis (EG/EoD) is significantly underdiagnosed and may be a common cause of moderate-to-severe gastrointestinal symptoms. Forty-five percent (45%, 181/405) of patients with chronic functional gastrointestinal symptoms who underwent upper endoscopy with biopsy met the diagnostic criteria for EG/EoD.
- Completed patient enrollment in Phase 3 EG/EoD and Phase 2/3 eosinophilic esophagitis (EoE) clinical trials of lirentelimab (AK002).

#### **Upcoming 2021 Milestones**

- Topline data from a randomized, double-blind, placebo-controlled Phase 3 study of lirentelimab in patients with EG/EoD expected in the fourth quarter of 2021.
- Topline data from a randomized, double-blind, placebo-controlled Phase 2/3 study of lirentelimab in patients with EoE expected in the fourth quarter of 2021.
- Initiation of a randomized, double-blind, placebo-controlled Phase 2/3 study of subcutaneous lirentelimab in patients with EG and/or EoD expected in the second half of 2021.
- Initiation of a Phase 2 study in a non-eosinophilic gastrointestinal disease in the second half of 2021.

#### **Second Quarter 2021 Financial Results**

Research and development expenses were \$41.0 million in the second quarter of 2021 as compared to \$28.3 million in the same period in 2020, an increase of \$12.7 million.

General and administrative expenses were \$16.2 million in the second quarter of 2021 as compared to \$12.1 million in the same period in 2020, an increase of \$4.1 million.

Allakos reported a net loss of \$57.2 million in the second quarter of 2021 as compared to \$39.3 million in the same period in 2020, an increase of \$17.9 million. Net loss per basic and diluted share was \$1.07 for the second quarter of 2021 compared to \$0.80 in the same period in 2020.

Allakos ended the second quarter of 2021 with \$559.7 million in cash, cash equivalents and marketable securities.

#### **About Allakos**

Allakos is a clinical stage biotechnology company developing antibodies that target immunomodulatory receptors present on immune effector cells involved in allergic, inflammatory, and proliferative diseases. The Company's lead antibody, lirentelimab (AK002), is being evaluated in a Phase 3 study in eosinophilic

gastritis (EG) and/or eosinophilic duodenitis (EoD) and a Phase 2/3 study in eosinophilic esophagitis (EoE). Lirentelimab targets Siglec-8, an inhibitory receptor selectively expressed on human mast cells and eosinophils. Inappropriately activated eosinophils and mast cells have been identified as key drivers in a number of severe diseases affecting the gastrointestinal tract, eyes, skin, lungs and other organs. Lirentelimab has been tested in multiple clinical studies. In these studies, lirentelimab eliminated blood and tissue eosinophils, inhibited mast cells and improved disease symptoms in patients with EG and/or EoD, EoE, mast cell gastrointestinal disease, severe allergic conjunctivitis, chronic urticaria and indolent systemic mastocytosis. For more information, please visit the Company's website at www.allakos.com.

#### **Forward-Looking Statements**

This press release contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995 as contained in Section 27A of the Securities Act of 1933, as amended, and Section 21E of the Securities Exchange Act of 1934, as amended. Such forward-looking statements include, but are not limited to, Allakos' progress and business plans, the expected timing of anticipated study results and plans relating to its future clinical trials. Such statements are subject to numerous important factors, risks and uncertainties that may cause actual events or results to differ materially from current expectations and beliefs, including but not limited to: Allakos' stages of clinical drug development; Allakos' ability to timely complete clinical trials for, and if approved, commercialize lirentelimab (AK002), its lead compound; Allakos' ability to obtain required regulatory approvals for its product candidates; uncertainties related to the enrollment of patients in its clinical trials; Allakos' ability to demonstrate sufficient safety and efficacy of its product candidates in its clinical trials; uncertainties related to the success of later-stage clinical trials, regardless of the outcomes of preclinical testing and early-stage trials; market acceptance of Allakos' product candidates; uncertainties related to the projections of the size of patient populations suffering from the diseases Allakos is targeting; Allakos' ability to advance additional product candidates beyond lirentelimab; Allakos' ability to obtain additional capital to finance its operations; and other important risk factors set forth in Allakos' most recent Annual Report on Form 10-K filed with the SEC on March 1, 2021, Quarterly Report on Form 10-Q filed with the SEC on August 9, 2021, and future reports to be filed with the SEC. These documents contain and identify important factors that could cause the actual results for Allakos to differ materially from those contained in Allakos' forward-looking statements. Any forward-looking statements contained in this press release speak only as of the date hereof, and Allakos specifically disclaims any obligation to update any forward-looking statement, except as required by law. These forward-looking statements should not be relied upon as representing Allakos' views as of any date subsequent to the date of this press release.

Source: Allakos Inc.

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# ALLAKOS INC. STATEMENTS OF OPERATIONS AND COMPREHENSIVE LOSS (in thousands, except per share data) (Unaudited)

	Three Months Ended			Six Months Ended				
	June 30,			June 30,				
		2021 2020		2021		2020		
Operating expenses								
Research and development	\$	40,985	\$	28,346	\$	79,900	\$	46,631
General and administrative		16,210		12,058		32,880		23,646
Total operating expenses		57,195		40,404		112,780		70,277
Loss from operations	<u> </u>	(57,195)		(40,404)		(112,780)		(70,277)
Interest income		103		1,284		233		3,273
Other expense, net		(117)		(172)		(220)		(112)
Net loss		(57,209)		(39,292)		(112,767)		(67,116)
Unrealized gain (loss) on marketable								
securities		(56)		(1,219)		24		650
Comprehensive loss	\$	(57,265)	\$	(40,511)	\$	(112,743)	\$	(66,466)
Net loss per common share:								
Basic and diluted	\$	(1.07)	\$	(0.80)	\$	(2.11)	\$	(1.38)
Weighted-average number of common								
shares outstanding:								
Basic and diluted		53,669		48,816	_	53,429	_	48,753

## ALLAKOS INC. CONDENSED BALANCE SHEETS (in thousands)

		June 30, 2021 (Unaudited)		December 31, 2020		
Assets						
Current assets:	Φ.	202.054	Φ.	205.455		
Cash and cash equivalents	\$	203,956	\$	207,177		
Investments in marketable securities		355,739		451,820		
Prepaid expenses and other current assets		16,363		10,270		
Total current assets		576,058		669,267		
Property and equipment, net		25,034		8,345		
Operating lease right-of-use assets		37,736		39,731		
Other long-term assets		16,119		2,275		
Total assets	\$	654,947	\$	719,618		
Liabilities and stockholders' equity						
Current liabilities:						
Accounts payable	\$	15,663	\$	13,960		
Accrued expenses and other current liabilities		18,335		8,490		
Total current liabilities		33,998		22,450		
Operating lease liabilities, net of current portion		49,963		42,773		
Total liabilities		83,961		65,223		
Stockholders' equity:						
Common stock		53		53		
Additional paid-in capital		1,026,632		997,298		
Accumulated other comprehensive gain		32		8		
Accumulated deficit		(455,731)		(342,964)		
Total stockholders' equity		570,986		654,395		
Total liabilities and stockholders' equity	\$	654,947	\$	719,618		