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): November 5, 2021

IMMUNOVANT, INC. (Exact name of Registrant as specified in its Charter)

			_
	Delaware	001-38906	83-2771572
	(State or other jurisdiction of incorporation or organization)	(Commission File Number)	(IRS Employer Identification No.)
	320 West 37th Street		
	New York, NY		10018
	(Address of principal executive offices)		(Zip Code)
	Registrant's telep	phone number, including area coo	de: (917) 580-3099
			_
Check the ap	ppropriate box below if the Form 8-K filing is intended to simulta	neously satisfy the filing obligation	of the registrant under any of the following provisions:
	Written communications pursuant to Rule 425 under the Secur	rities Act (17 CFR 230.425)	
	Soliciting material pursuant to Rule 14a-12 under the Exchang	ge Act (17 CFR 240.14a-12)	
	Pre-commencement communications pursuant to Rule 14d-2(b	b) under the Exchange Act (17 CFR	240.14d-2(b))
	Pre-commencement communications pursuant to Rule 13e-4(c	e) under the Exchange Act (17 CFR	240.13e-4(c))
Securities re	gistered pursuant to Section 12(b) of the Act:		
	Title of each class	Trading Symbol(s)	Name of each exchange on which registered
	Common Stock, \$0.0001 par value per share	IMVT	The Nasdaq Stock Market LLC
	check mark whether the registrant is an emerging growth company ct of 1934 (§240.12b-2 of this chapter).	y as defined in Rule 405 of the Secr	urities Act of 1933 (§230.405 of this chapter) or Rule 12b-2 of the Securities
Emerging gr	rowth company		
	ng growth company, indicate by check mark if the registrant has ovided pursuant to Section 13(a) of the Exchange Act. \Box	elected not to use the extended trans	sition period for complying with any new or revised financial accounting

Item 2.02 Results of Operations and Financial Condition.

On November 5, 2021, Immunovant, Inc., or the Company, issued a press release announcing its financial results for its fiscal second quarter and six months ended September 30, 2021. A copy of the press release is furnished as Exhibit 99.1 to this Current Report on Form 8-K and is incorporated herein by reference.

The information contained in this Item 2.02, including Exhibit 99.1, shall not be deemed "filed" for purposes of Section 18 of the Securities Exchange Act of 1934, as amended, or subject to the liabilities of that section or Sections 11 and 12(a)(2) of the Securities Act of 1933, as amended. The information in this Current Report shall not be incorporated by reference in any filing with the U.S. Securities and Exchange Commission made by the Company, whether made before or after the date hereof, regardless of any general incorporation language in such filing.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits

Exhibit No.	Description
99.1	Press Release dated November 5, 2021
104	Cover Page Interactive Data File (embedded within the 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.

Immunovant, Inc.

By: /s/ Eva Renee Barnett

Eva Renee Barnett Chief Financial Officer

Date: November 5, 2021

Immunovant Reports Financial Results for the Quarter Ended September 30, 2021 Company Ended the Quarter With Cash of Approximately \$559 Million

NEW YORK, November 5, 2021 (GLOBE NEWSWIRE) mmunovant, Inc. (Nasdaq: IMVT), a clinical-stage biopharmaceutical company focused on enabling normal lives for people with autoimmune diseases, today reported financial results for its fiscal first quarter ended September 30, 2021. Immunovant ended the quarter with approximately \$559 million in cash.

Financial Highlights for Fiscal Second Quarter Ended September 30, 2021:

R&D Expenses: Research and development expenses were \$21.4 million for the three months ended September 30, 2021, compared to \$12.0 million for the three months ended September 30, 2020. The year-over-year increase primarily reflected higher contract manufacturing costs and personnel-related expenses (including stock-based compensation), reflecting investment spending to support our strategic objectives as we prepare to re-initiate our clinical activities. These increases were partially offset by lower program-specific clinical trial activities due to the continued voluntary pause.

G&A Expenses: General and administrative expenses were \$16.3 million for the three months ended September 30, 2021, compared to \$9.0 million for the three months ended September 30, 2020. The year-over-year increase was primarily due to financial advisory fees, legal and other professional costs and higher personnel-related expenses (including stock-based compensation).

Net Loss: Net loss was \$37.7 million (\$0.35 per common share) for the three months ended September 30, 2021, compared to \$20.8 million (\$0.25 per common share) for the three months ended September 30, 2020. Net loss for the three months ended September 30, 2021 and 2020 included \$8.4 million and \$3.4 million, respectively, related to non-cash stock-based compensation expense.

Common Stock: As of September 30, 2021, there were 114,998,871 shares of common stock issued and outstanding.

Financial Highlights for Fiscal Six Months Ended September 30, 2021:

R&D Expenses: Research and development expenses were \$40.1 million for the six months ended September 30, 2021, compared to \$28.9 million for the six months ended September 30, 2020. The year-over-year increase primarily reflected higher personnel-related expenses (including stock-based compensation), increases in clinical studies and clinical research and higher contract manufacturing costs, reflecting investment spending to support our strategic objectives as we prepare to re-initiate our clinical activities. These increases were partially offset by lower program-specific clinical trial activities due to the continued voluntary pause.

G&A Expenses: General and administrative expenses were \$27.5 million for the six months ended September 30, 2021, compared to \$18.7 million for the six months ended September 30, 2020. The year-over-year increase was primarily due to financial advisory fees, legal and other professional costs and higher personnel-related expenses (including stock-based compensation).

Net Loss: Net loss was \$68.2 million (\$0.66 per common share) for the six months ended September 30, 2021, compared to \$47.5 million (\$0.61 per common share) for the six months ended September 30, 2020 Net loss for the six months ended September 30, 2021 and 2020 included \$12.2 million and \$7.3 million, respectively, related to non-cash stock-based compensation expense.

About Immunovant, Inc.

Immunovant, Inc. is a clinical-stage biopharmaceutical company focused on enabling normal lives for patients with autoimmune diseases. Immunovant is developing IMVT-1401 ("batoclimab"), a novel, fully human anti-FcRn monoclonal antibody, as a subcutaneous injection for the treatment of autoimmune diseases mediated by pathogenic IgG antibodies.

Forward-Looking Statements

This press release contains forward-looking statements for the purposes of the safe harbor provisions under The Private Securities Litigation Reform Act of 1995 and other federal securities laws. The use of words such as "may," "might," "would," "should," "expect," "believe," "estimate," and other similar expressions are intended to identify forward-looking statements. Such forward looking statements include Immunovant's plan to develop batoclimab across a broad range of autoimmune indications. All forward-looking statements are based on estimates and assumptions by Immunovant's management that, although Immunovant believes to be reasonable, are inherently uncertain. All forward-looking statements are subject to risks and uncertainties that may cause actual results to differ materially from those that Immunovant expected. Such risks and uncertainties include, among others: initial results or other preliminary analyses or results of early clinical trials may not be predictive final trial results or of the results of later clinical trials; the timing and availability of data from clinical trials; the timing of discussions with regulatory agencies, as well as regulatory submissions and potential approvals; the continued development of Immunovant's product candidate, including the timing of the commencement of additional clinical trials and resumption of current trials; Immunovant's scientific approach, clinical trial design, indication selection and general development progress; future clinical trials may not confirm any safety, potency or other product characteristics described or assumed in this

press release; any product candidate that Immunovant develops may not progress through clinical development or receive required regulatory approvals within expected timelines or at all; Immunovant's product candidate may not be beneficial to patients, or even if approved by regulatory authorities, successfully commercialized; the potential impact of the ongoing COVID-19 pandemic on Immunovant's clinical development plans and timelines; Immunovant's business is heavily dependent on the successful development, regulatory approval and commercialization of its sole product candidate, batoclimab; Immunovant is at an early stage in development of batoclimab; and Immunovant will require additional capital to fund its operations and advance batoclimab through clinical development. These and other risks and uncertainties are more fully described in Immunovant's periodic and other reports filed with the Securities and Exchange Commission (SEC), including in the section titled "Risk Factors" in Immunovant's most recent Annual Report on Form 10-K, its Form 10-Q to be filed with the SEC on November 5, 2021, and Immunovant's subsequent filings with the SEC. Any forward-looking statement speaks only as of the date on which it was made. Immunovant undertakes no obligation to publicly update or revise any forward-looking statement, whether as a result of new information, future events or otherwise.

IMMUNOVANT, INC.

Condensed Consolidated Statements of Operations

(Unaudited, in thousands, except share and per share data)

	Three Months Ended September 30,			Six Months Ended September 30,				
		2021		2020		2021		2020
Operating expenses:								
Research and development	\$	21,361	\$	11,976	\$	40,066	\$	28,898
General and administrative		16,289		8,998		27,469		18,662
Total operating expenses		37,650		20,974		67,535		47,560
Other expense (income), net		84		(225)		711		(151)
Loss before (benefit) provision for income taxes		(37,734)		(20,749)		(68,246)		(47,409)
(Benefit) provision for income taxes		(31)		40		(72)		88
Net loss	\$	(37,703)	\$	(20,789)	\$	(68,174)	\$	(47,497)
Net loss per common share – basic and diluted	\$	(0.35)	\$	(0.25)	\$	(0.66)	\$	(0.61)
Weighted-average common shares outstanding – basic and diluted		109,078,427		84,353,438		103,558,036		77,623,132

IMMUNOVANT, INC.

Condensed Consolidated Balance Sheets

(Unaudited, in thousands, except share and per share data)

	September 30, 2021		March 31, 2021		
Assets					
Current assets:					
Cash	\$	558,952	\$	400,146	
Prepaid expenses and other current assets		4,840		8,860	
Total current assets		563,792		409,006	
Operating lease right-of-use assets		2,731		3,282	
Property and equipment, net		209		201	
Total assets	\$	566,732	\$	412,489	
Liabilities and Stockholders' Equity					
Current liabilities:					
Accounts payable	\$	4,243	\$	2,432	
Accrued expenses		23,058		15,160	
Current portion of operating lease liabilities		1,082		1,179	
Total current liabilities		28,383		18,771	
Operating lease liabilities, net of current portion, and other noncurrent liabilities		1,952		2,238	
Total liabilities		30,335		21,009	
Commitments and contingencies					
Stockholders' equity:					
Series A preferred stock, par value \$0.0001 per share, 10,000 shares authorized, issued and outstanding at September 30, 2021 and March 31, 2021		_		_	
Preferred stock, par value \$0.0001 per share, 10,000,000 shares authorized, no shares issued and outstanding at September 30, 2021 and March 31, 2021		_		_	
Common stock, par value \$0.0001 per share, 500,000,000 shares authorized, 114,998,871 shares issued and outstanding at September 30, 2021 and 500,000,000 shares authorized, 97,971,243 shares issued and outstanding at March 31, 2021					
		12		10	
Additional paid-in capital		802,774		590,425	
Accumulated other comprehensive income (loss)		442		(298)	
Accumulated deficit		(266,831)		(198,657)	
Total stockholders' equity		536,397		391,480	
Total liabilities and stockholders' equity	Ś	566,732	Ś	412,489	

Contact:

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