#### **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): January 14, 2025

# 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 telephone number, including area code: (917) 580-3099								
Check the	appropriate box below if the Form 8-K filing is intended to simult	aneously satisfy the filing obligation of	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))							
	Pre-commencement communications pursuant to Rule 13e-4(c) under the Exchange Act (17 CFR 240.13e-4(c))							
Securities 1	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, \$0.0001 par value per share	IMVT	The Nasdaq Stock Market LLC					
	check mark whether the registrant is an emerging growth comparator of 1934 (§240.12b-2 of this chapter).	ny as defined in Rule 405 of the Secur	ities Act of 1933 (§230.405 of this chapter) or Rule 12b-2 of the Securities					
Emerging g	growth company □							
	ging growth company, indicate by check mark if the registrant has provided pursuant to Section 13(a) of the Exchange Act. $\Box$	elected not to use the extended transit	tion period for complying with any new or revised financial accounting					

#### Item 7.01. Regulation FD Disclosure.

Immunovant, Inc. (the "Company") will provide a corporate overview for investors with a new corporate presentation at the 43rd Annual J.P. Morgan Healthcare Conference on January 14, 2025. A copy of the presentation is attached hereto as Exhibit 99.1 and is incorporated herein by reference.

The information furnished under this Item 7.01, 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 Item 7.01, including Exhibit 99.1, shall not be deemed incorporated by reference into any other 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

Presentation, dated January 14, 2025.

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99.1

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: January 14, 2025

Exhibit 99.1



# Targeted science, + Tailored solutions +

for people with autoimmune disease



2025 J.P. Morgan Healthcare Conference

January 14, 2025 <sub>+</sub>



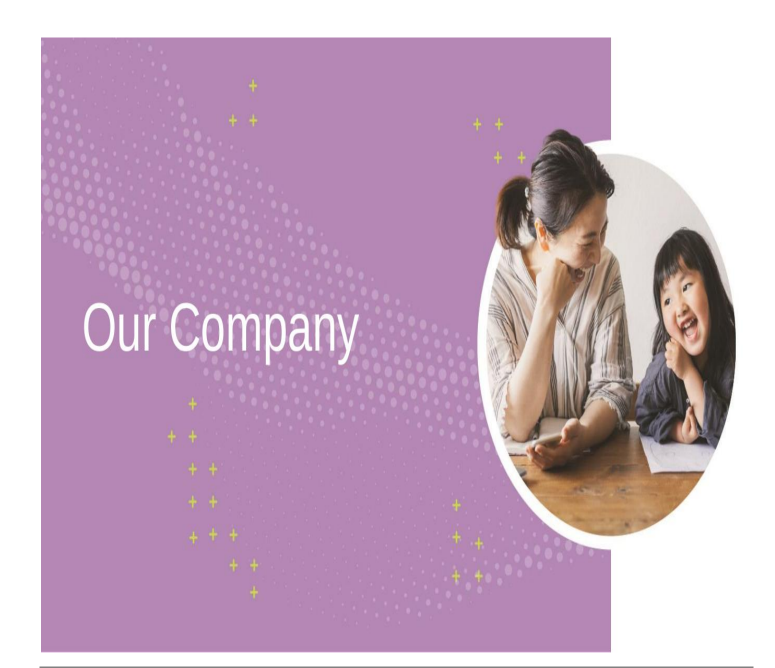
## Forward-looking statements

This presentation 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 "can," "may," "might," "will," "would," "should," "expect," "believe," "estimate," "design," "plan," "intend," "anticipate," and other similar expressions are intended to identify forward-looking statements. Such forward looking statements include Immunovant's expectations regarding patient enrollment, timing, design, and results of clinical trials of its product candidates and indication selections; Immunovant's plan to develop IMVT-1402 and batoclimab across a broad range of autoimmune indications; expectations with respect to these planned clinical trials including the number and timing of (a) trials Immunovant expects to initiate, (b) FDA clearance with respect to IND applications, and (c) potential pivotal or registrational programs and clinical trials of IMVT-1402; the size and growth of the potential markets for Immunovant's product candidates and indication selections, including any estimated market opportunities; Immunovant's plan to explore in subsequent study periods follow-on treatment with alternative dosing regimens; Immunovant's beliefs regarding the potential benefits of IMVT-1402's and batoclimab's unique product attributes and first-in-class or best-in-class potential, as applicable; Immunovant's anticipated strategic reprioritization from batoclimab to IMVT-1402; and whether, if approved, IMVT-1402 or batcolimab will be successfully distributed, marketed or commercialized. 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 of final trial results or of the results of later clinical trials; results of animal studies may not be predictive of results in humans; 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 candidates, including the timing of the commencement of additional clinical 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 presentation; 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 candidates may not be beneficial to patients, or even if approved by regulatory authorities, successfully commercialized; the effect of global factors such as geopolitical tensions and adverse macroeconomic conditions on Immunovant's business operations and supply chains, including its clinical development plans and timelines; Immunovant's business is heavily dependent on the successful development, regulatory approval and commercialization of batoclimab and IMVT-1402; Immunovant is in various stages of clinical development for IMVT-1402 and batoclimab; and Immunovant will require additional capital to fund its operations and advance IMVT-1402 and 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 Quarterly Report on Form 10-Q for the quarter ended September 30, 2024, filed with the SEC on November 7, 2024, 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® and IMMUNOVANT® are registered trademarks of Immunovant Sciences GmbH. All other trademarks, trade names, service marks, and copyrights appearing in this presentation are the property of their respective owners. Dates used in this presentation refer to the applicable calendar year unless otherwise noted.



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## Our vision:

# Normal lives for people with autoimmune disease

#### What we do:

We are developing targeted therapies that are designed to address the complex and variable needs of people with autoimmune diseases.



Love Trailblazing



Bolder, Faster



All Voices





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# 2024: Many milestones achieved supporting lead asset IMVT-1402

Graves' POC observed greater benefit with deeper IgG reduction

5 INDs cleared for lead asset, IMVT-1402

Initiated IMVT-1402 pivotal trials in Graves' Disease & ACPA+ D2T RA¹

Unprecedented speed of starting pivotal trials with autoinjector²

MG trial completed enrollment with batoclimab

Meaningfully strengthened balance sheet



## Our focus:

Pursue a broad anti-FcRn strategy based on potential best-in-class profile of IMVT-1402 targeting autoantibody-driven diseases



development and commercialization experience across the C-suite and senior leaders

protection for IMVT-1402 to 2043<sup>1</sup>

Issued U.S. claims cover composition of matter, method of use, and methods for manufacturing

equivalents totaled approximately \$473M as of September 30, 2024

Gross proceeds of \$450M from PIPE January 2025

regulatory approval of efgartigimod and rozanolixizumab

best-in-class profile

5 INDs cleared including GD, ACPA+ D2T RA, MG and CIDP<sup>2</sup>

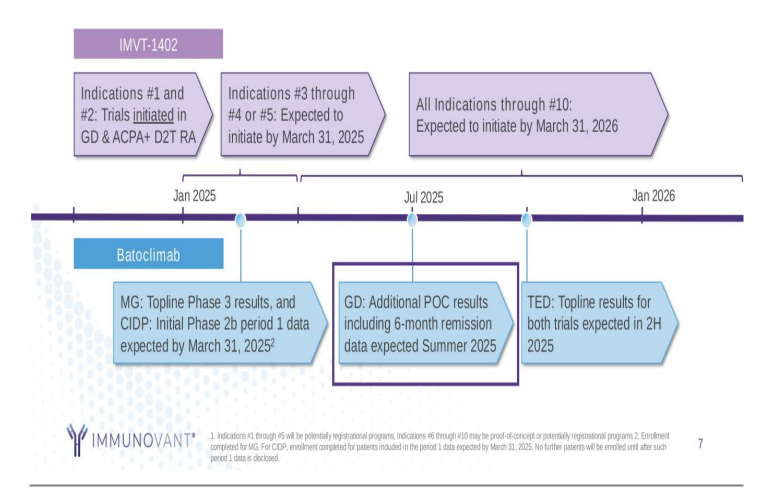
with 23 indications announced or in development across the anti-FcRn class<sup>3</sup>



- 1. Not including any potential patent term extension
  2. Anti-citrullinated protein autoantibody positive (ACPA+), Difficult-to-Treat Rheumatoid Arthritis (D2T RA), Myasthenia Gravis (MG), Chronic Inflammatory Demyelinating
- Indications announced or in development with anti-FcRn assets by Immunovant, argenx, Johnson & Johnson, and UCB

## Multiple near-term milestones for enhanced value creation

On track to initiate 4-5 potentially registrational programs for IMVT-1402 by March 31, 2025 and trials in a total of 10 indications by March 31, 2026<sup>1</sup>



# Graves' data demonstrates potentially transformational results in patients uncontrolled on ATDs with greater response driven by deeper IgG lowering

Phase 2 batoclimab proof of concept data



# Graves' US market-sizing analyses confirm high unmet need with ~330K prevalent patients relapsed, uncontrolled, or intolerant to ATDs

- Conservative Inovalon claims analysis¹ yields <u>~880K prevalent</u> **Graves' Disease patients,** including <u>~330K prevalent</u> ATD relapsed patients choosing not to pursue ablation
- Conservative Inovalon claims analysis<sup>2</sup> yields <u>~65K annual incident</u> **Graves' Disease patients**, including <u>~20K annual incident</u> second line uncontrolled / intolerant patients
- Deep dive endocrinologist survey of 140 healthcare providers treating Graves' Disease patients indicates ~25-30% of patients are relapsed, uncontrolled, or intolerant to ATDs
- Real-world chart audit of 1,120 Graves' Disease patients treated by surveyed endocrinologists indicates ~25-30% of patients are relapsed, uncontrolled, or intolerant to ATDs
- Patient survey of 100 diagnosed Graves' Disease patients indicates ~25-30% of patients are relapsed, uncontrolled, or intolerant to ATDs



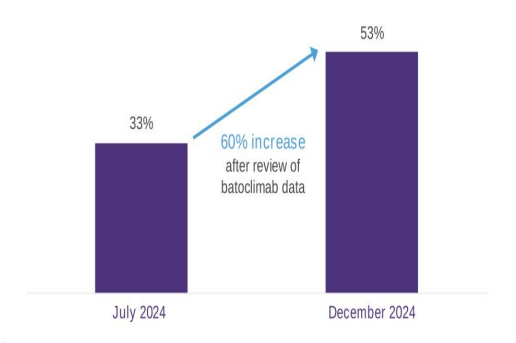
. Roivant Claims Analysis – 2022 prevalent patient population based on a two-year lookback for diagnosis

2. Roivant Claims Analysis – 2021 incident patient population based on a five-year lookback to define the incident population

Note: See Immunovant, Inc. Graves' Disease Program Update Deck dated September 9, 2024, available at Immunovant.com

# Unmet medical need in Graves' disease was rated higher by thyroid specialists after exposure to batoclimab data

### Percent of ATD-treated GD patients needing alternative medical therapy

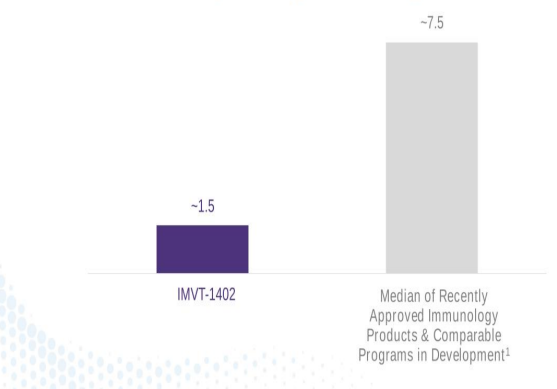




Source: Graves' Disease HCP Quantitative Survey (n=152 Endocrinologists) by Immunovant, July 202:

# Unprecedented speed of starting pivotal trials with an autoinjector<sup>1</sup>







Includes benralizumab, bimekizumab, dupilumab, gefurulimab, guselkumab, ixekizumab, mepolizumab, mirkizumab, risankizumab, sarilumab, secukinumab, sonetokimab, tralokinumab, tezepelumab, VRDN-003 and zilucoptan

Measured from start of first-in-human study to start of first trial with autoinjector; studies include but are not limited to efficacy, safety, bioequivalence, self-injection, device

# IMVT-1402 starting pivotal trials with intended commercial formulation and device: YpsoMate® autoinjector

Leveraging market-proven, user-friendly technology to meet patient needs

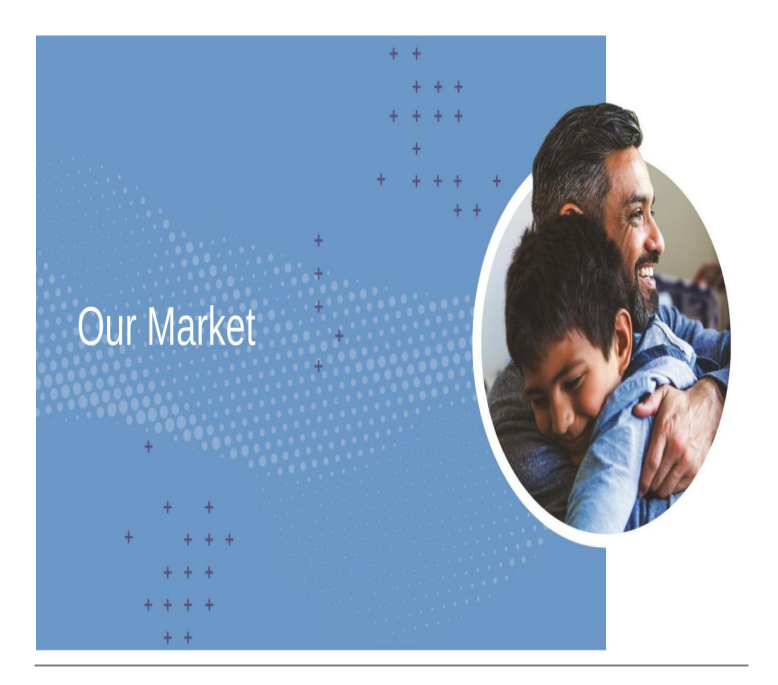


Established autoinjector with multiple approved products

- · Automated, simple, subcutaneous injection
- · Hidden needle shield
- Provides both visual and audio feedback



"Ypsomate" autoinjector used in ADBRY®, COSENTYX®, AJOVY®, NUCALA®, FASENRA®, TEZSPIRE® "Ypsomate" is a registered trademark of Ypsomed AG.



# 2024: Many positive developments for the FcRn inhibitor class



Positive data in new indications<sup>1</sup>



Approval in new indication<sup>2</sup>



Mixed results from other modalities<sup>3</sup>



Growing KOL enthusiasm for earlier line anti-FcRn use

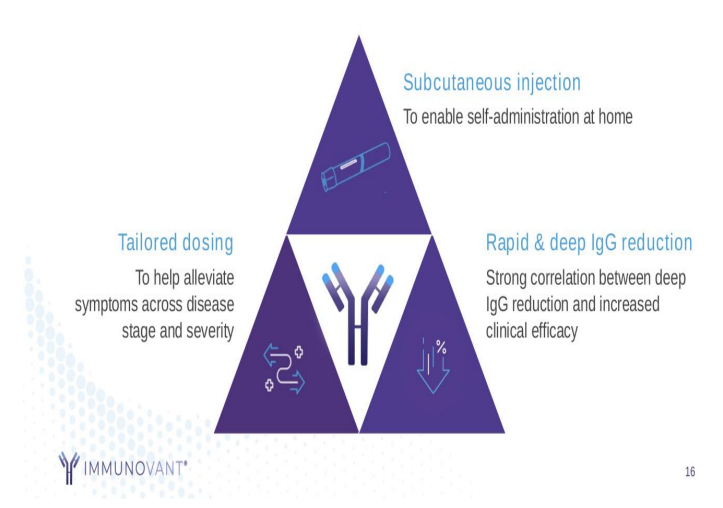
# Ever-growing conviction in anti-FcRn as a uniquely exciting class



- https://us.argenx.com/news/2024/argenx-advances-clinical-development-efgartigimod-primary-sjogrens-disease; https://www.janssen.com/late-breaking-results-show-nipocalimab-significantly-improves-sjogrens-disease-activity-phase-2
   Wygart Hytrulo approval for CIDP; https://www.fda.gov/drugs/news-events-human-drugs//da-approves-treatment-chronic-inflammatory-demyelinating-polyneuropathy-cidp-
- Wygart Hytrulo approval for CIDP; https://www.lda.gov/drugs/news-events-human-drugs/fda-approves-treatment-chronic-inflammatory-demyelinating-polyneuropathy-ddj adults
- https://www.chugai-pharm.co.jp/english/news/detail/20240321150000\_1059.html

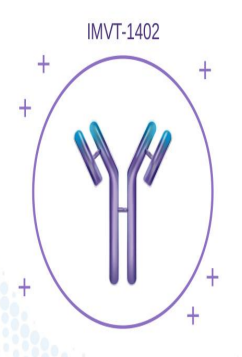
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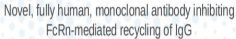
# Our differentiated value proposition: Three potentially unique attributes to address unmet patient needs



## Our lead asset:

# IMVT-1402 has a combination of potentially best-in-class attributes not seen with other anti-FcRns







Deep IgG Lowering Phase 1 data suggests deep dosedependent IgG lowering



Favorable Analyte Profile Phase 1 data supports a favorable analyte profile with no or minimal effect on albumin and LDL



Convenient Administration Delivered via market-proven, user-friendly autoinjector



Compelling Patent Protection Issued U.S. patent covers composition of matter, method of use and methods for manufacturing to 2043<sup>1</sup>



Not including any potential patent term extension

# An exciting 2025





# 2025: Exciting year ahead

- MG and CIDP data (CYQ1) and TED data (CYH2) designed to reinforce correlation of greater efficacy with deeper IgG reduction
- Additional data from Graves' POC including 6-month remission data designed to further articulate potential for IMVT-1402 in Graves'
- Potentially registrational trials enrolling in GD, ACPA+ D2T RA, MG, CIDP and soon to unveiled 5<sup>th</sup> indication
  - Additional studies (including POCs) to be announced for IMVT-1402, all with autoinjector
- O5 Studies initiated in 10 indications by March 31, 2026



## Multiple near-term milestones for enhanced value creation

On track to initiate 4-5 potentially registrational programs for IMVT-1402 by March 31, 2025 and trials in a total of 10 indications by March 31, 2026<sup>1</sup>

