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

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**FORM 8-K**

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**CURRENT REPORT**  
**Pursuant to Section 13 or 15(d)**  
**of the Securities Exchange Act of 1934**

**Date of Report (Date of earliest event reported): September 7, 2022**

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

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**Delaware**  
(State or other jurisdiction of incorporation or organization)

**001-38906**  
(Commission File Number)

**83-2771572**  
(IRS Employer Identification No.)

**320 West 37th Street**  
**New York, NY**  
(Address of principal executive offices)

**10018**  
(Zip Code)

**Registrant's telephone number, including area code: (917) 580-3099**

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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:

- ☐ 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:

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

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 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. ☐

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**Item 7.01 Regulation FD Disclosure.**

On September 7, 2022, Immunovant, Inc., or the Company, will host a pre-announced corporate update webcast and conference call. A copy of the presentation to be used during the webcast 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, or the Exchange Act, or subject to the liabilities of that section or Sections 11 and 12(a)(2) of the Securities Act of 1933, or the Securities Act. 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 Exchange Commission, or the SEC, made by the Company, whether made before or after the date hereof, regardless of any general incorporation language in such filing.

**Item 8.01 Other Events.**

On September 7, 2022, the Company issued a press release announcing plans to initiate a Pivotal Phase 2b trial in Chronic Inflammatory Demyelinating Polyneuropathy (CIDP) in the second half of calendar year 2022, with initial results from open-label period 1 expected in the first half of calendar year 2024, and plans to initiate a Phase 2 trial in Graves’ Disease in early 2023, with initial results expected in the second half of calendar year 2023. The Company also plans to finalize its trial design in Warm Autoimmune Hemolytic Anemia following interactions with regulators later in 2022.

A copy of this press release is attached hereto as Exhibit 99.2. The information contained in this Item 8.01 is hereby incorporated by reference into the Company’s Registration Statement on Form S-3 (File No. 333-251865).

**Item 9.01 Financial Statements and Exhibits.**

(d) Exhibits

<u>Exhibit No.</u>	<u>Description</u>
99.1	<a href="#">Investor Presentation dated September 7, 2022.</a>
99.2	<a href="#">Press release dated September 7, 2022.</a>
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, 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: September 7, 2022



# Immunovant Announces Plans to Study Batoclimab in Two New Indications



*Committed to enabling normal lives for people with autoimmune disease*

September 7, 2022



# 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," and other similar expressions are intended to identify forward-looking statements. Such forward looking statements include Immunovant's plan to initiate a Phase 2b clinical trial for batoclimab in Chronic Inflammatory Demyelinating Polyneuropathy in the second half of calendar year 2022 with initial results from open-label period 1 expected in the first half of calendar year 2024; Immunovant's plan to initiate a Phase 2 clinical trial for batoclimab in Graves' Disease in early 2023 with initial results expected in the second half of calendar year 2023; Immunovant's plan to report topline data from its Phase 3 trial for batoclimab in Myasthenia Gravis in the second half of calendar year 2024; Immunovant's plan to initiate two Phase 3 clinical trials for batoclimab in Thyroid Eye Disease in the second half of calendar year 2022 with expected topline data readouts in the first half of calendar year 2025; Immunovant's plan to finalize its trial design in Warm Autoimmune Hemolytic Anemia following expected interactions with regulators later in calendar year 2022; Immunovant's plan to develop batoclimab across a broad range of autoimmune indications; expectations with respect to the safety and monitoring plan and size of the safety database for these planned clinical trials; the timing of discussions with regulatory agencies; the size and growth of the potential markets for Immunovant's product candidates and indication selections; Immunovant's plan to explore in subsequent study periods follow-on treatment with alternative dosing regimens; Immunovant's expectations regarding patient enrollment, timing, the design and results of clinical trials of its product candidates and indication selections; Immunovant's beliefs regarding its cash runway, and the potential benefits of batoclimab's unique product attributes. 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 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 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, geopolitical tensions, and adverse macroeconomic conditions 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 filed with the SEC on August 5, 2022, 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.


*All 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.*



# Q&A housekeeping

You may submit questions through the Q&A function at the bottom of the webcast player. Please see the image here for reference.

Starts Sep 7th at 8:00 AM EDT



Immunovant

Please fill out the form below to ask our presenters live questions.

Name	Company	Question	( * - mandatory fields)
<input type="text"/>	<input type="text"/>	<input type="text"/>	<input type="button" value="Submit Your Question"/>



# Today's agenda

## Immunovant and batoclimab

### *Introduction*

- Pete Salzmann, MD, CEO Immunovant

## Chronic Inflammatory Demyelinating Polyneuropathy (CIDP)

### *Differentiated development program to optimize effect size*

- Jonathan Katz<sup>1</sup>, MD, Director Neuromuscular Clinic, California Pacific Medical Center
- Todd Levine<sup>2</sup>, Medical Director, Neurology, Honor Health Scottsdale, Arizona
- Pete Salzmann, MD, CEO Immunovant

## Graves' Disease

### *First in class development program in high unmet need population*

- George Kahaly<sup>3</sup>, MD, PhD, Johannes Gutenberg University Medical Center
- Pete Salzmann, MD, CEO Immunovant

## Closing

### *Summary of milestones and catalysts*

- Pete Salzmann, MD, CEO Immunovant

## Q&A



**Financial Disclosures:** 1. Consultant for Argenx, Grifols, MT Pharma, Biogen, Amylyx, Alexion, PTC Therapeutics, Calico, UCB; 2. Eledon, Speakers' bureau for Grifols. Financial interests in CND Life Sciences and CRL. Consult for InCircle; 3. The Johannes Gutenberg University (JGU) Medical Center, Mainz, Germany (academic institution of George J. Kahaly, MD, PhD) has received research-associated funding from the JGU Medical Faculty, AdvanceCor (Germany), Apitope (Belgium), Berlin-Chemie (Germany), Byondis (The Netherlands), GlycoEra (Switzerland), Horizon (USA), Immunovant (USA), ISAR (Germany), Medimics (USA), Merck (Germany), Novartis (USA), Quidel (USA), River Vision (USA), and Roche (Switzerland). GJK consults for GlycoEra, Immunovant, ISAR, Medimics, Merck, Novartis, Quidel, & VasaraGen (USA).

# Pipeline expansion: Chronic Inflammatory Demyelinating Polyneuropathy (CIDP) & Graves' Disease

01

CIDP is an exciting opportunity for the anti-FcRn class. We believe that our **uniquely optimized trial** could position batoclimab with best-in-class efficacy and first-to-market simple SC

02

Graves' Disease is a first-in-class opportunity in an autoantibody driven condition with a **straightforward Phase 2** approach and a **high unmet need** in a large subset of patients insufficiently responding even at maximally tolerated doses of current standard of care

03

Data catalysts for new indications complement pivotal data from MG and TED programs. Expect **cadence of data every half year** starting in the second half of 2023

04

Immunovant has \$427M in cash with cash runway into 2025<sup>1,2</sup>



SC = Subcutaneous administration; MG = Myasthenia Gravis; TED = Thyroid Eye Disease

1. As of June 30, 2022, per most recent Quarterly Report on Form 10-Q filed with the SEC on August 5, 2022
2. The assumptions upon which we have based our estimates, including expenditures relating to planned or potential clinical trials, are routinely evaluated and may be subject to change

For Investor Audiences Only **5**



# Immunovant and Batoclimab

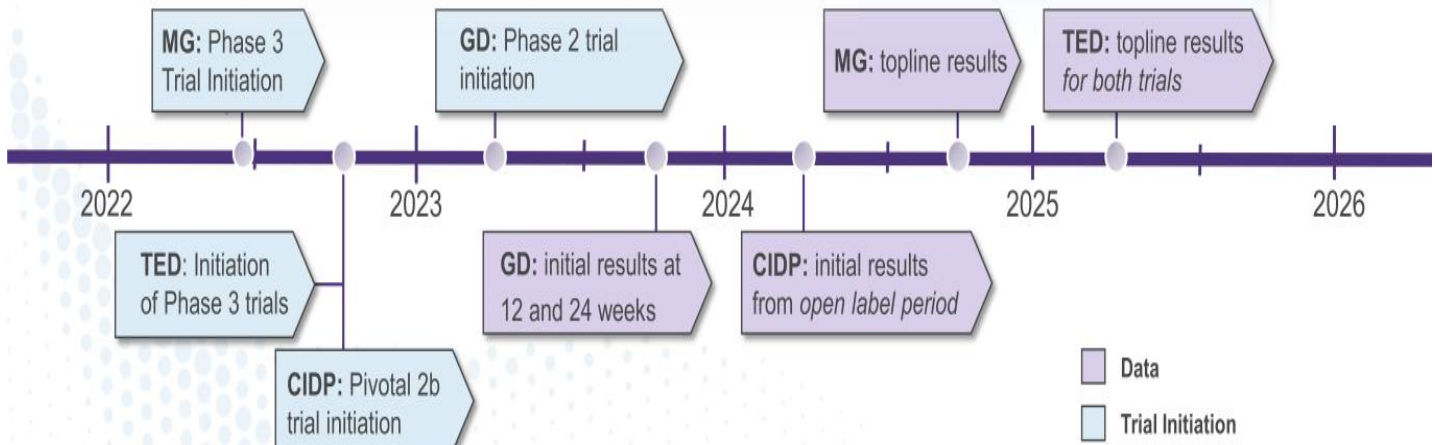
*Pete Salzmann, MD  
Chief Executive Officer*



# Pursuing a broad development program with batoclimab

## Planning for regular cadence of data across indications

Target Indication	Stage of Development
Myasthenia Gravis (MG)	Pivotal Phase 3
Thyroid Eye Disease (TED)	Pivotal Phase 3
Chronic Inflammatory Demyelinating Polyneuropathy (CIDP)	Pivotal Phase 2b*
Graves' Disease (GD)	Phase 2
Warm Autoimmune Hemolytic Anemia (WAIHA)	Phase 2**



\*Registrational package for CIDP may include 1 or 2 pivotal trials depending on a variety of factors

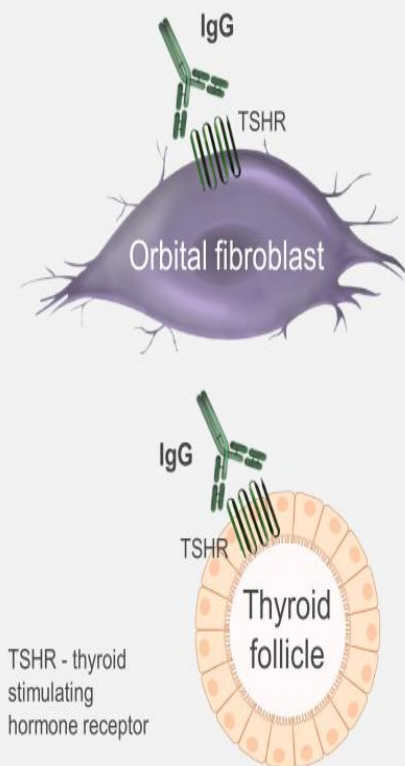
\*\*WAIHA design to be finalized after FDA interaction expected in 2H of calendar 2022

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# IgG antibodies play a role in autoimmune disease pathogenesis

- In many autoimmune diseases, IgG antibodies develop that bind to normal tissues<sup>1</sup>
- Some IgG autoantibodies trigger harmful immune responses resulting in autoimmune symptoms and tissue damage
- Some IgG autoantibodies bind cell-surface receptors that may be activated
- **Disease severity may correlate with quantity of pathogenic IgG**

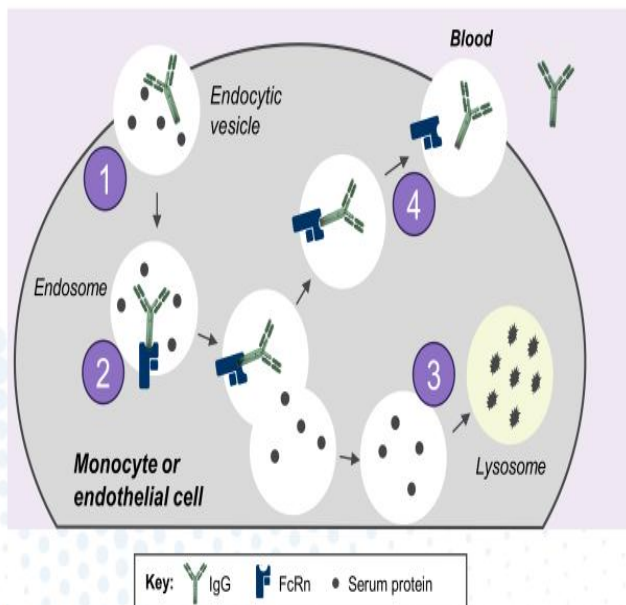
## Normal tissues recognized by IgG autoantibodies in Thyroid Eye Disease<sup>1</sup>



## FcRn promotes recycling of IgG antibodies

- FcRn extends the half-life of IgG autoantibodies in circulation exacerbating their autoimmune effects
- FcRn expressed in a variety of cells

### FcRn maintains levels of IgG in circulation by preventing IgG degradation



#### FcRn Mechanism of Action

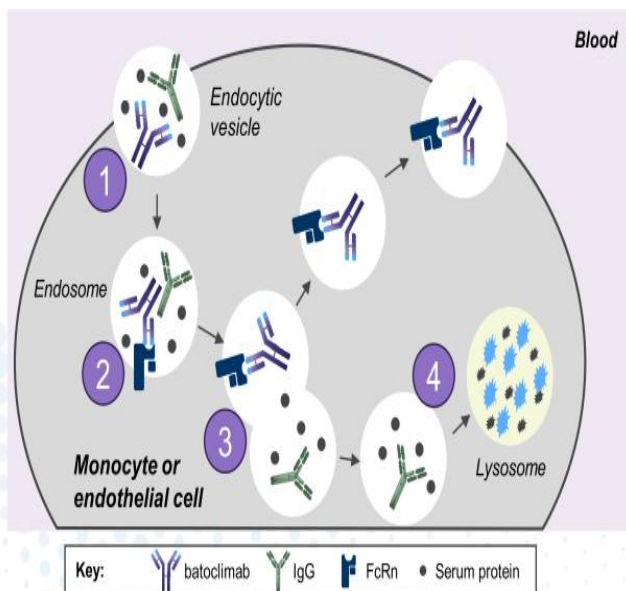
1. IgG is taken up into cells in endocytic vesicle
2. FcRn-IgG complexes are sorted from unbound proteins
3. Unbound proteins are trafficked to lysosome for degradation
4. IgG is recycled back into circulation



# Batoclimab inhibits FcRn, promoting IgG degradation

- Batoclimab binds to FcRn and reduces the recycling of IgG antibodies
- As a result, IgG is increasingly delivered to lysosomes for degradation
- Relative to older, broad-spectrum immunosuppressants, FcRn inhibitors deliver a more targeted approach to immunomodulation

## Batoclimab removes pathogenic antibodies by binding to FcRn and promoting IgG degradation



### Batoclimab Mechanism of Action

1. IgG and batoclimab are taken up into cells in endocytic vesicles
2. Batoclimab binds to FcRn in endosomes
3. FcRn-batoclimab complexes are sorted from unbound proteins
4. Non-receptor bound IgGs are degraded in lysosomes

# Chronic Inflammatory Demyelinating Polyneuropathy (CIDP)

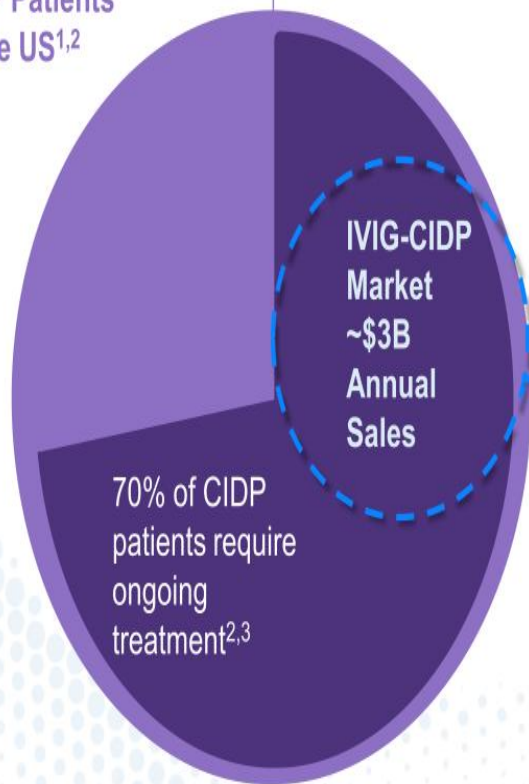
New Indication Announcement 2022





# CIDP represents an exciting opportunity

16,000 Total  
CIDP Patients  
in the US<sup>1,2</sup>



Batoclimab has the potential to enhance  
CIDP-IVIG market with a favorable risk-  
benefit-tolerability profile

- CIDP represents 22% of total IVIG market by volume: ~\$3B in global annual sales for IVIG in CIDP<sup>4</sup>
- Existing CIDP treatments can produce side effects and have logistical challenges
- Target population: Active CIDP patients



Sources: 1. Broers M, et al (2019) Incidence and prevalence of CIDP: a systematic review and meta-analysis. *Neuroepidemiology* 52(3-4):161-172; 2. Querol L, et al. Systematic literature review of burden of illness in chronic inflammatory demyelinating polyneuropathy (CIDP). *J Neurol* 268, 3706-3716 (2021); 3. Kuijwaard K, Bos-Eysen ME, Blomkwist-Markens PH et al (2009) Recurrences, vaccinations and long-term symptoms in GBS and CIDP. *J Periph Nerv Syst* 14(4):310-315. <https://doi.org/10.1111/j.1529-8027.2009.00243.x>; 4. CSL Behring R&D Investor Briefing, 2021.

Fireside chat on  
**CIDP**



**Pete Salzmann, MD**  
Chief Executive Officer,  
Immunovant



**Todd Levine, MD**  
Medical Director, Neurology  
Department, Honor Health  
Scottsdale, Arizona



**Jonathan Katz, MD**  
Director Neuromuscular  
Clinic, California Pacific  
Medical Center

# Key Takeaways from CIDP Discussion

01

We believe CIDP is a very important disease in neurology and an exciting potential indication for the anti-FcRn class as current therapies (IVIG, PLEX and steroids) are effective, but have **significant side effects and logistical limitations** (IVIG & PLEX).

02

Diagnosis of CIDP is challenging and must be done carefully. **Algorithms may improve diagnostic accuracy**, which is paramount for clinical trial success.

03

Clinical trial design is also critical to **maximize ability to demonstrate effect size**. Patients entering CIDP trials after IVIG withdrawal are essentially untreated, whereas patients entering CIDP trials after steroid reduction remain partially treated and this may blunt the effect size observed in trials that include these patients within the primary analysis group.

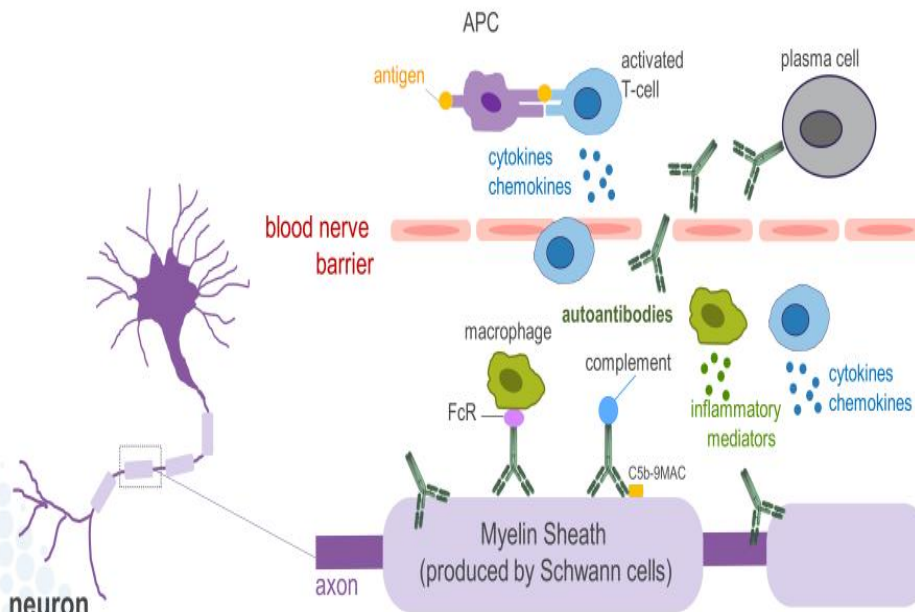
04

CIDP is a chronic, symptomatic condition for many patients and therefore an **effective treatment** that could be administered via **a simple subcutaneous injection** would represent **a meaningful improvement for patients** with CIDP. We believe CIDP is ripe for disruption with the right mechanism, the right asset and the right trial design.



IVIG = intravenous immunoglobulin

# Response to IVIG and Plasma Exchange creates strong rationale for potential benefit of anti-FcRn mechanism even in cases without known auto-antibody



CIPD is characterized by predominant demyelination of motor and sensory nerves. Although the root cause of CIPD is unknown, significant evidence suggests that the disorder(s) are immunologically mediated.<sup>1</sup>

Multiple immune mechanisms including cellular (macrophages), humoral and complement pathways contribute to the pathogenesis of CIPD.<sup>2,3</sup>

anti-myelinated peripheral nerve IgG in 30-40% patients<sup>1</sup>



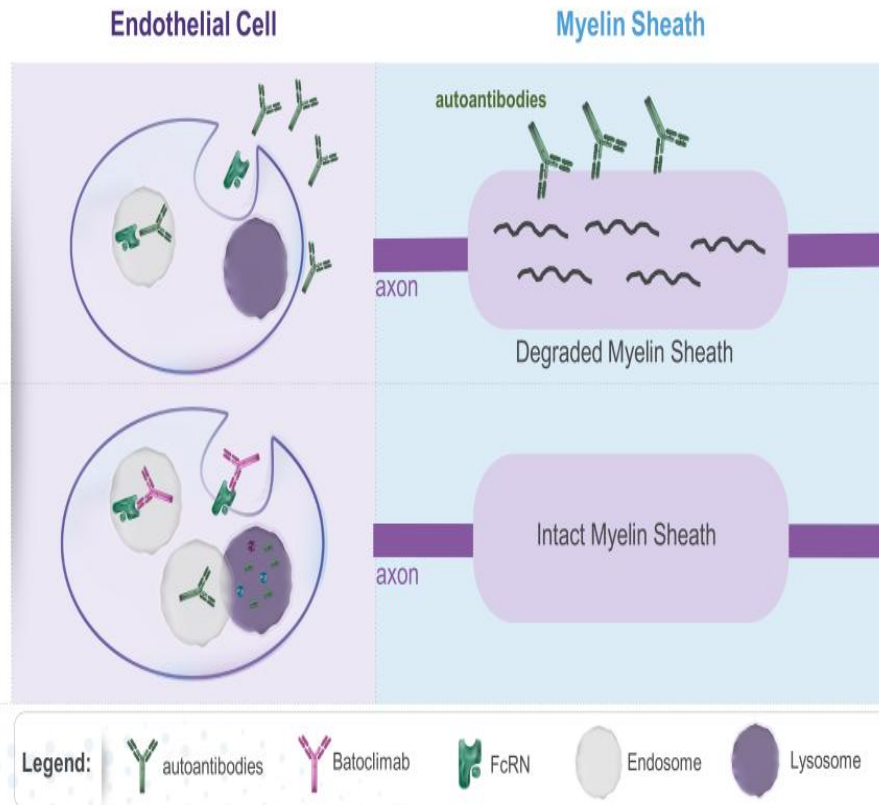
Sources: 1. Mathey EK, Park SB, Hughes RAC, et al Chronic inflammatory demyelinating polyradiculoneuropathy: from pathology to phenotype Journal of Neurology, Neurosurgery & Psychiatry 2015; 2. Koike H, Katsuno M. Pathophysiology of Chronic Inflammatory Demyelinating Polyneuropathy: Insights into Classification and Therapeutic Strategy. Neurol Ther. 2020 Dec;9(2):213-227. doi: 10.1007/s40120-020-00190-8. Epub 2020 May 14; 3. Querol LA, Hartung HP, Lewis RA, van Doorn PA, Hammond TR, Atassi N, Alonso-Alonso M, Dalakas MC. The Role of the Complement System in Chronic Inflammatory Demyelinating Polyneuropathy: Implications for Complement-Targeted Therapies. Neurotherapeutics. Apr 2022.



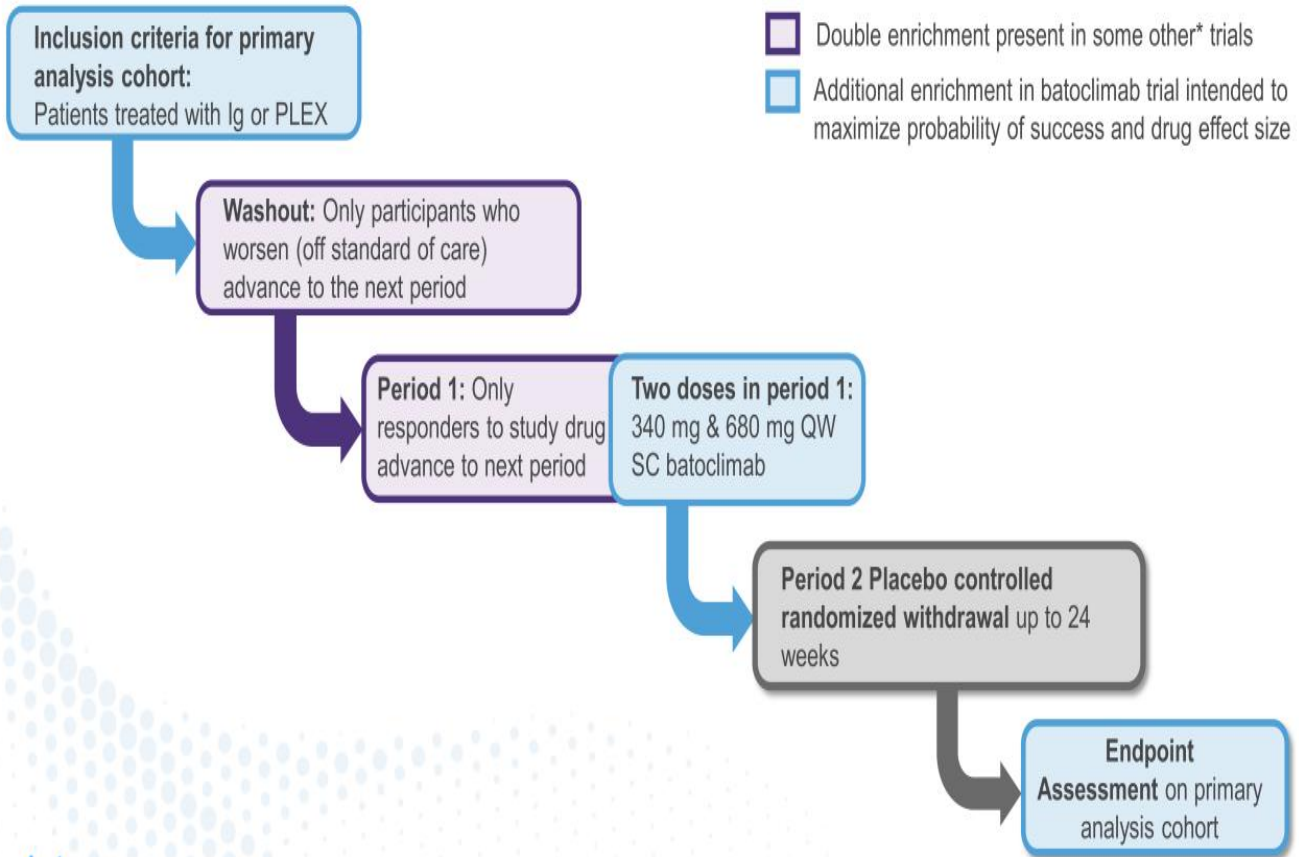
## Anti-FcRn mechanism of action degrades IgG, potentially protecting the myelin sheath from pathogenic IgG autoantibody attack in CIDP

**In the absence of batoclimab,** FcRn binds to the IgG autoantibodies, inhibiting their degradation and returning them into circulation. IgGs may attack the myelin resulting in myelin degradation and neuropathy

**With batoclimab,** FcRn is blocked from binding to IgG autoantibodies, which are then transported to the lysosome for degradation, decreasing their levels in circulation and potentially reducing pathogenesis



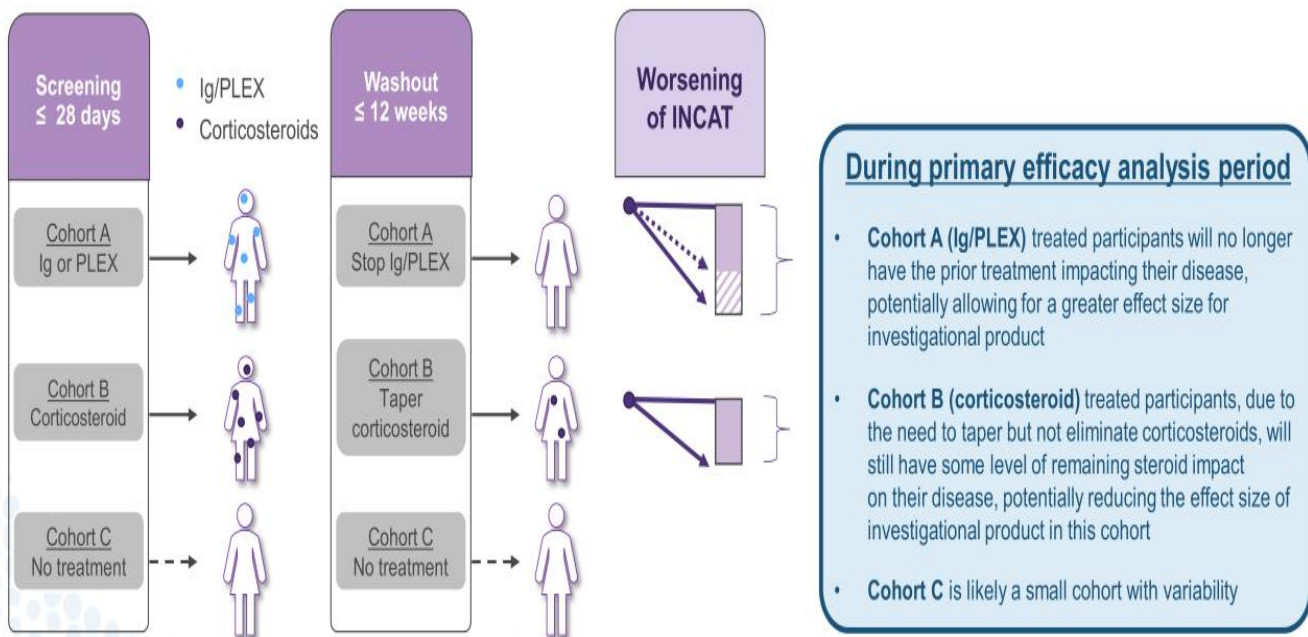
## Batoclimab trial in CIDP builds on learning from recent trials with a novel triple enrichment strategy



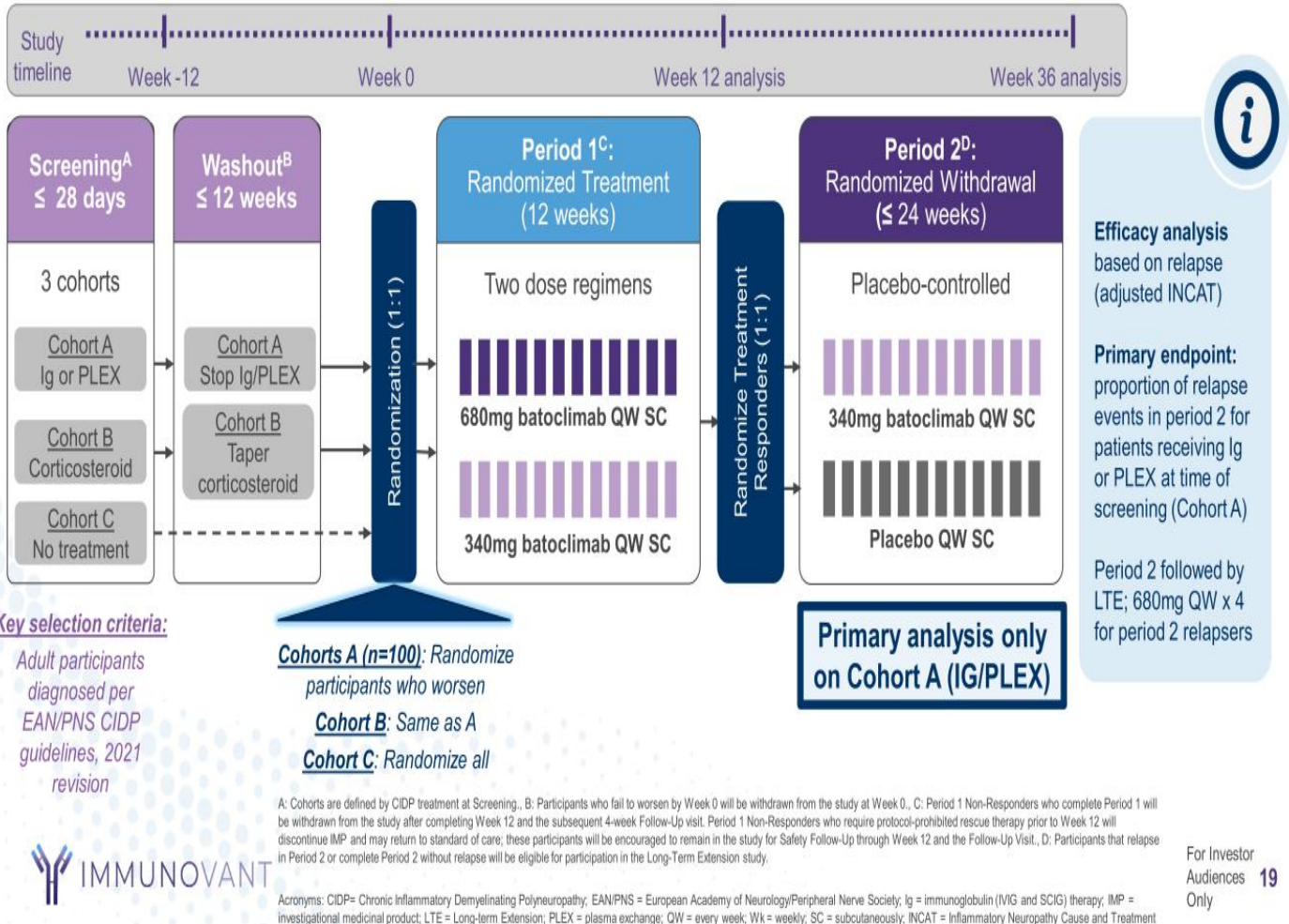
Note: \*Other trials includes anti-FcRn's in development by Argenx and Janssen; Source: Argenx and Janssen public disclosures  
Acronyms: Ig = immunoglobulin (IVIg and SCIG) therapy; PLEX = plasma exchange; QW = Dosed weekly; SC = Subcutaneous administration



Restricting the primary analysis to Cohort A patients previously on Ig/PLEX maximizes the number of participants fully off prior therapy for the primary endpoint and may improve separation from placebo



# CIDP pivotal Phase 2b trial design intended to enable development of potentially best-in-anti-FcRn-class chronic therapy for CIDP



## Our development approach applies key learnings from historical and ongoing CIDP trials to address challenges unique to CIDP

Trial risks	Mitigations	Mitigation included in other anti-FcRn Trials*	Mitigation included in IMVT trial
Disease heterogeneity and challenging diagnosis	<b>Diagnostic algorithm</b>	X	✓
Mix of active and inactive disease among those enrolled creates risk of low relapse rate in the placebo arm, thereby shrinking potential effect size for the investigational product	<b>Double enrichment:</b> 1. Subjects must worsen after withdrawal / taper of standard of care (e.g. IVIG/SCIG, PLEX, or steroids) on study entry, AND 2. Subjects must then improve on open label investigational product	Not All**	✓
Patients enrolled in placebo arm of trial may not have demonstrated initial response to investigational product		Not All**	✓
Steroids are a common standard of care outside the US and often can't be fully tapered, weakening double enrichment	<b>Third enrichment:</b> Primary endpoint on IVIG/SCIG/Plex cohort only to <b>maximize the potential effect size</b>	X	✓
Lack of dose exploration	Data on <b>multiple doses</b> in "Period 1" of trial will inform future development strategy	X	✓
Single large trial limits flexibility to optimize product label and differentiation	Smaller, initial pivotal trial that can be adjusted or complemented with a second smaller pivotal trial to optimize product label and differentiation based on new data	X	✓

**Staged approach (with or without additional trial if that is required for registration) has the potential to deliver a differentiated product label with a larger effect size**



Notes: \*Other anti-FcRn trials in CIDP include efgartigimod, nupocalmab, and rozanolixizumab. \*\*clinical trial designs for efgartigimod in CIDP and nupocalmab in CIDP include double enrichment in trial design. Rozanolixizumab ph2 trial in CIDP did not include double enrichment.

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## Immunovant pursuing a differentiated approach to developing batoclimab as a chronic treatment for CIDP

1

**CIDP is an exciting indication that is ripe for disruption**

Given disease complexity, trial design is critical

2

**Our pivotal study is optimized versus historical and current studies**

To improve probability of success and effect size, and include multiple doses for optimal differentiation

3

**Batoclimab has potential best-in-class efficacy and could be first simple SC**

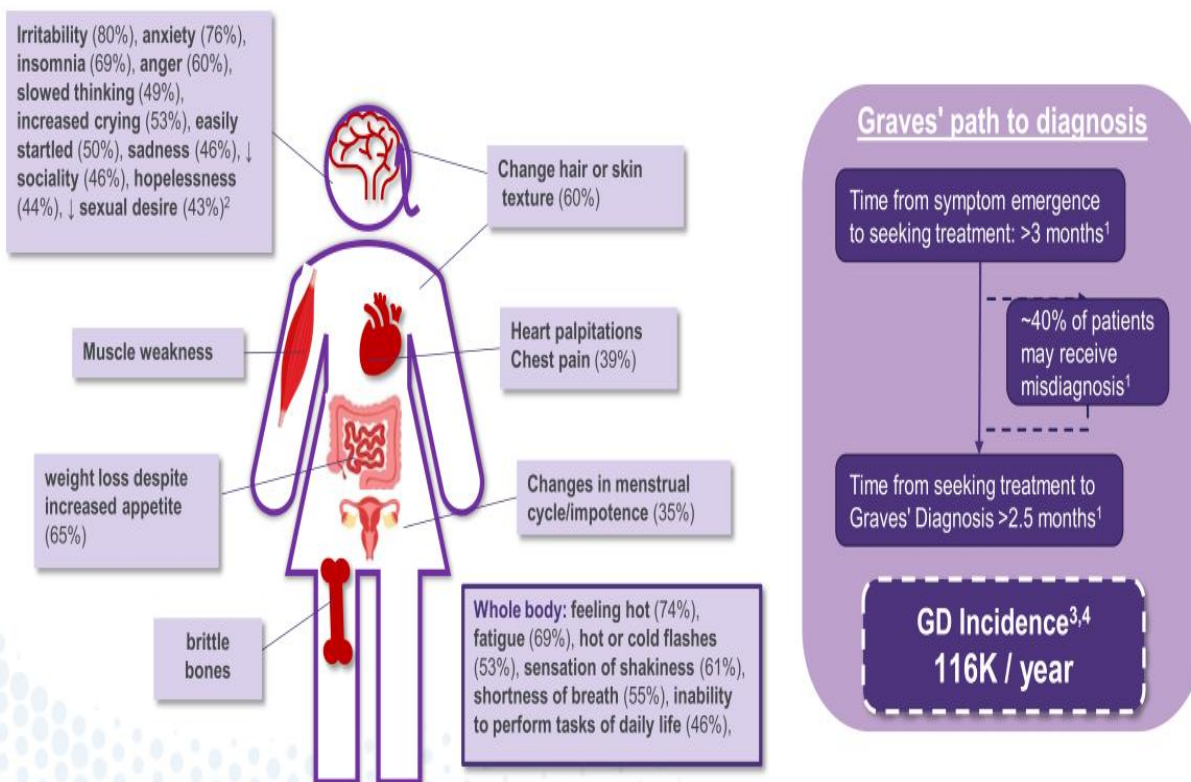
Representing meaningful innovation for patients with this chronic disease

# Graves' Disease

New Indication Announcement 2022



# Systemic Graves' Disease symptoms impact many organ systems and leave many patients with substantial symptoms **despite maximal tolerated medical therapy**

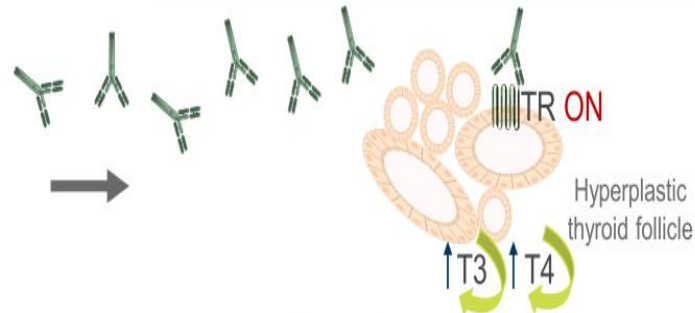
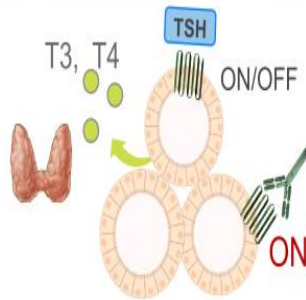




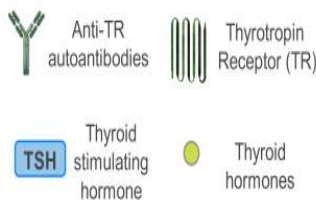
# Pathogenic anti-Thyrotropin Receptor (TR) autoantibodies turn 'ON' the TR signaling pathways leading to thyroid hyperplasia and systemic<sup>1-7</sup> Graves' disease

anti-TR autoantibodies (TR Ab)  
stimulate Thyrotropin Receptor on thyroid follicles

TR stimulation results in thyroid follicle hyperplasia and increased release of thyroid hormones

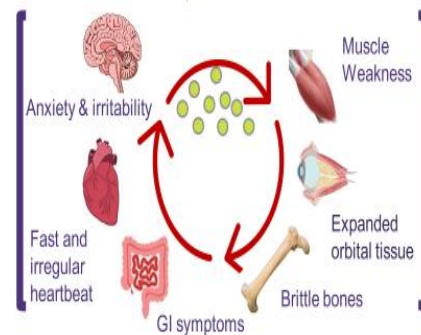


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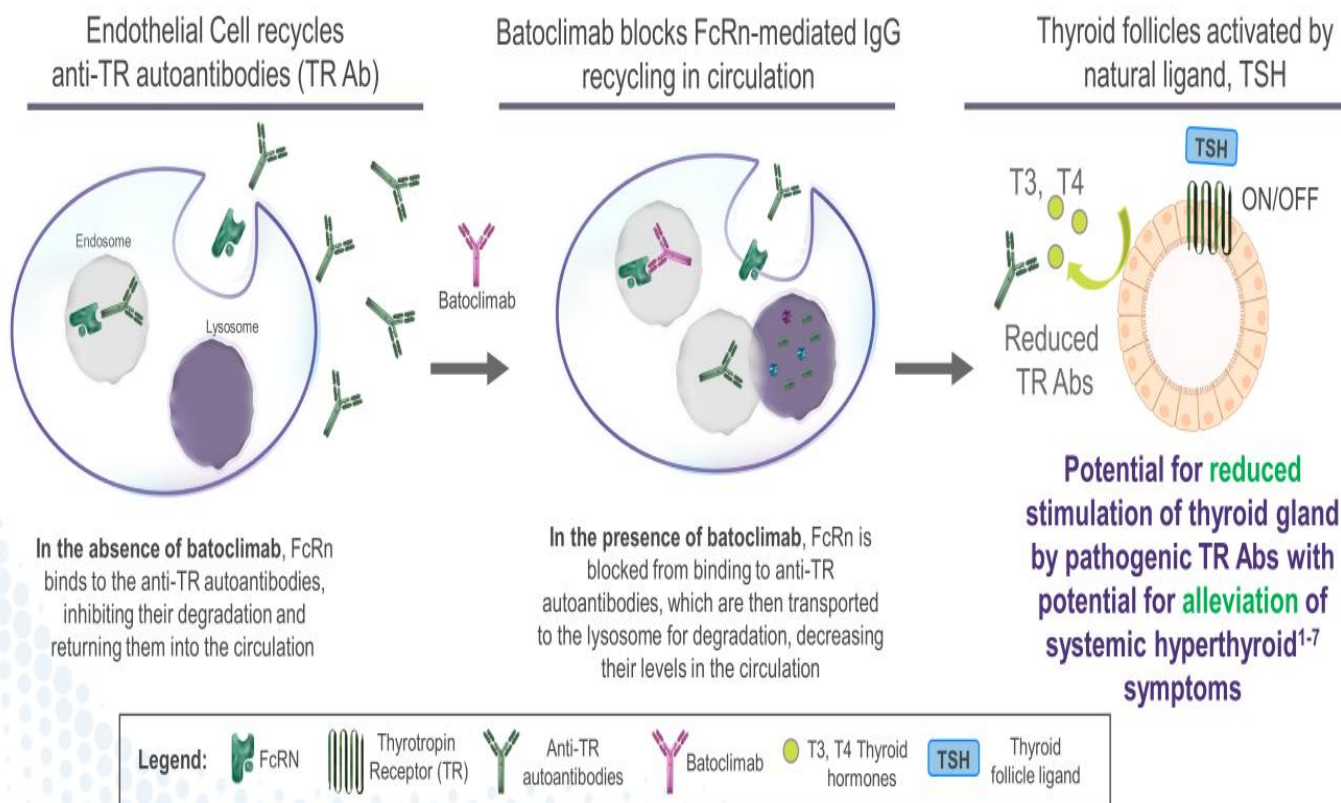
## Hyperthyroidism leads to multi-system<sup>1-7</sup> Graves' symptoms

Heart, skeletal muscle, skin, bones, eyes, liver, brain and reproductive and GI systems may be affected



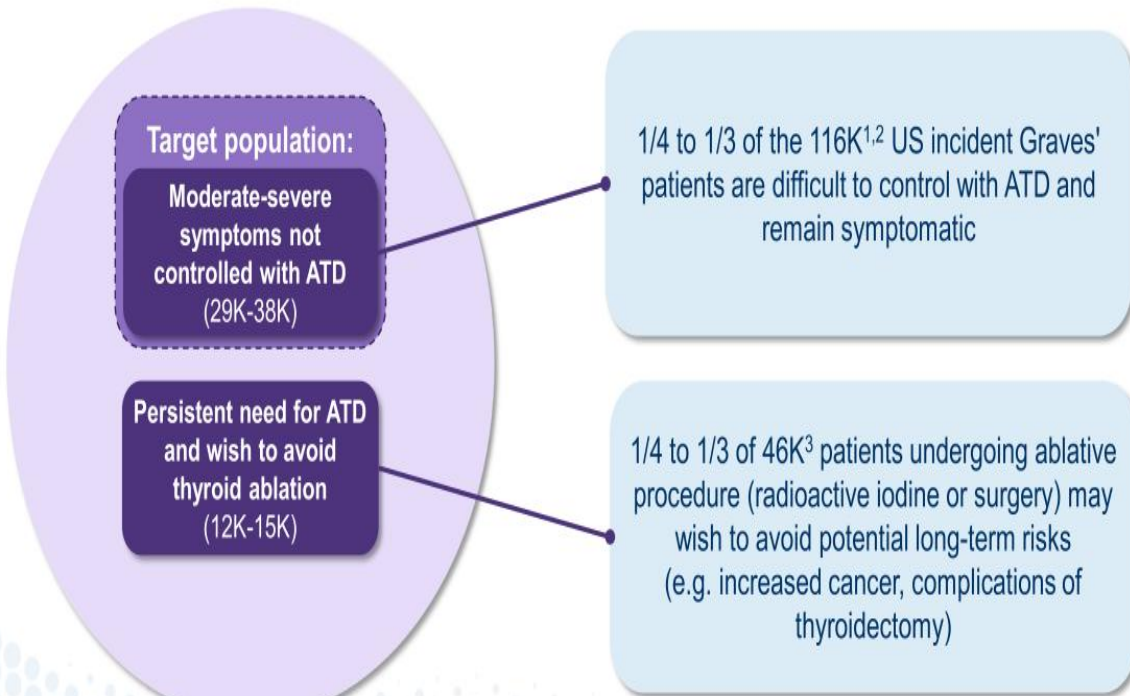
Sources: 1. Girgis CM, Champion BL, Wall JR. Current concepts in Graves' disease. *Ther Adv Endocrinol Metab.* 2011 Jun;2(3):135-44; 2. Gawalko M, Balsam P, Lodziński P, Grabowski M, Krzowski B, Opolski G, Kosiuk J. Cardiac Arrhythmias in Autoimmune Diseases. *Circ J.* 2020 Apr 24; 3. Fukao A, Takamatsu J, Arishima T, Tanaka M, Kawai T, Okamoto Y, Miyauchi A, Imagawa A. Graves' disease and mental disorders. *J Clin Transl Endocrinol.* 2019 Oct 11; 4. Kubota S, Amino N, Matsumoto Y, Ikeda N, Morita S, Kudo T, et al. (2008) Serial changes in liver function tests in patients with thyrotoxicosis induced by Graves' disease and painless thyroiditis. *Thyroid* 18, 283-287 5. Maser C, Toset A, Roman S. Gastrointestinal manifestations of endocrine disease. *World J Gastroenterol.* 2006 May 28; 6. Dhanwal DK. Thyroid disorders and bone mineral metabolism. *Indian J Endocrinol Metab.* 2011 Jul; 7. Sethi PP, Parchani A, Pathania M. Respiratory Muscle Weakness in Thyrotoxic Periodic Palsy. A Lesson to Remember. *Ann Neurosci.* 2021 Jul; 8. George's Kahaly GJ (2010) The thyrocyte-fibrocyte link: closing the loop in the pathogenesis of Graves' disease? *J Clin Endocrinol Metab* 95(1):62-65; 9. Chen H, Mester T, Raychaudhuri N, Kauh CY, Gupta S, Smith TJ, et al. Teprotumumab, an IGF-1R blocking monoclonal antibody inhibits TSH and IGF-1 action in fibrocytes. *J Clin Endocrinol Metab.* 2014 Sep

# Inhibiting FcRn to foster degradation of circulating autoantibodies may reduce thyroid hyperactivity and alleviate systemic<sup>1-7</sup> symptoms



Many patients with Graves' Disease remain symptomatic despite maximally tolerated anti-thyroid medications; others would avoid ablation if possible

A Total Addressable Incidence Population of 41K – 53K per year (US) beyond ATD





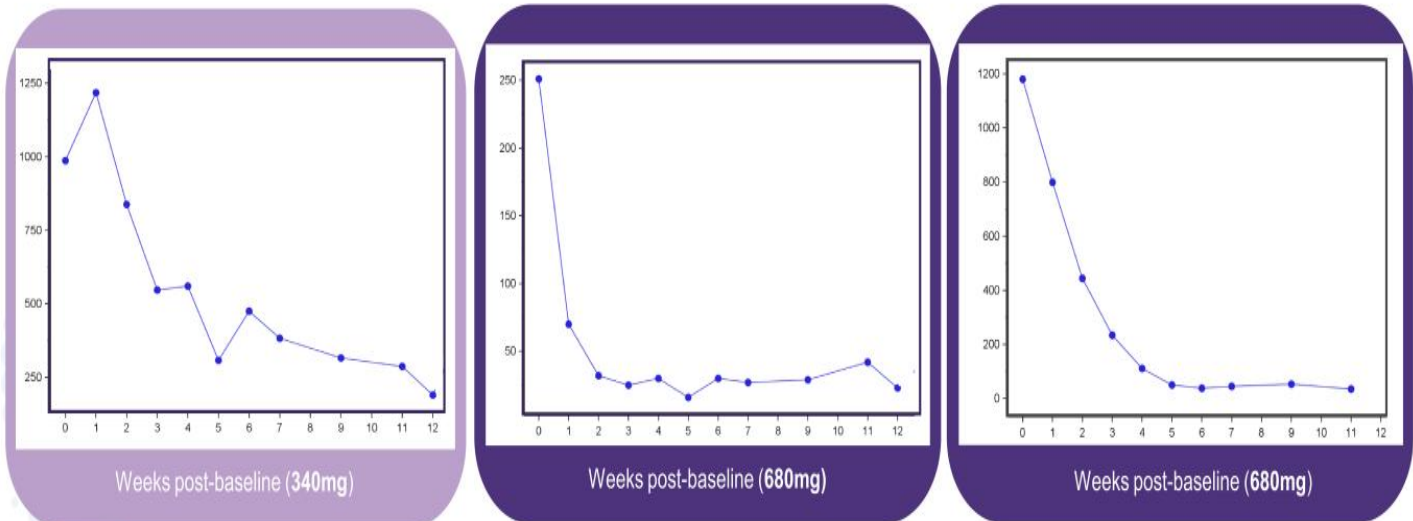
Many patients with Graves' Disease remain symptomatic despite maximally tolerated anti-thyroid medications and ablation can be problematic

	Safety			Tolerability		
SoC Treatments	Risk of liver damage	Risk of secondary cancers	Risk of low blood cell counts	Invasive	Rash/Itching	Hypothyroidism risk and fatigue
Anti-Thyroid Medicines	✓	X	✓	X	✓	✓
Radio Iodine	X	✓	X	X	X	✓
Surgery	X	X	X	✓*	X	✓

\*Surgical risks include laryngeal nerve damage, hypoparathyroidism and bleeding

## Observed reductions in stimulatory anti-TSHR antibodies with batoclimab from paused TED Phase 2b trial provide encouraging anecdotes for batoclimab in Graves' Disease

Level of Stimulatory Anti-TSHR Antibody (SRR%<sup>1</sup>) in TED Phase 2b study



While not representative of the population as a whole, three example patient responses to batoclimab in Phase 2b study in TED showed reductions in stimulatory anti-TSHR antibodies at both 340mg and 680mg doses of batoclimab



Source: Data on File, Immunovant, Inc.  
<sup>1</sup>SRR is the "Sample to Reference Ratio". This cell-based assay readout is the ratio of the sample signal to that of a reference control, expressed as %.  
A value less than 140 is considered negative for stimulatory antibody; a value greater than or equal, positive for stimulatory antibody.  
The efficacy of batoclimab and clinical outcomes were deemed inconclusive, in part, because the study was terminated early.

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Fireside chat on

# Graves' Disease



**Pete Salzmann, MD**  
Chief Executive Officer, Immunovant



**George J. Kahaly, MD, PhD**  
Professor of Medicine and Endocrinology/Metabolism  
Johannes Gutenberg University (JGU) Medical Center  
Department of Medicine I  
ORPHAN Disease Center for Graves' Orbitopathy  
and Autoimmune Polyendocrinopathy



# Key Takeaways from Graves' Disease Discussion

01

Standard of care for Graves' Disease is often limited by safety and tolerability concerns, leaving many patients needing additional efficacy to control their thyroid hormone levels and symptoms

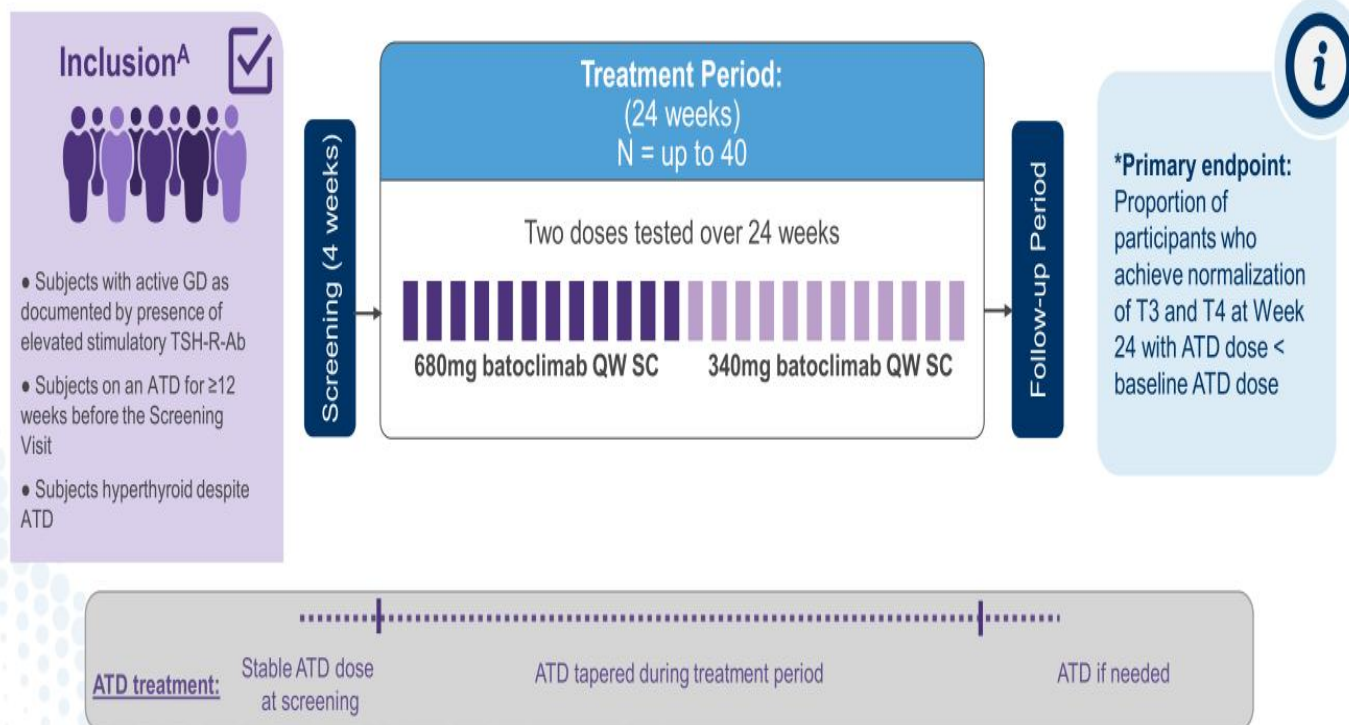
02

Diverse and bothersome symptoms primarily relate to elevated thyroid hormone levels (measured as increased T3 and T4) are a direct result of overstimulation of the thyroid gland by anti-TSHR auto-antibodies

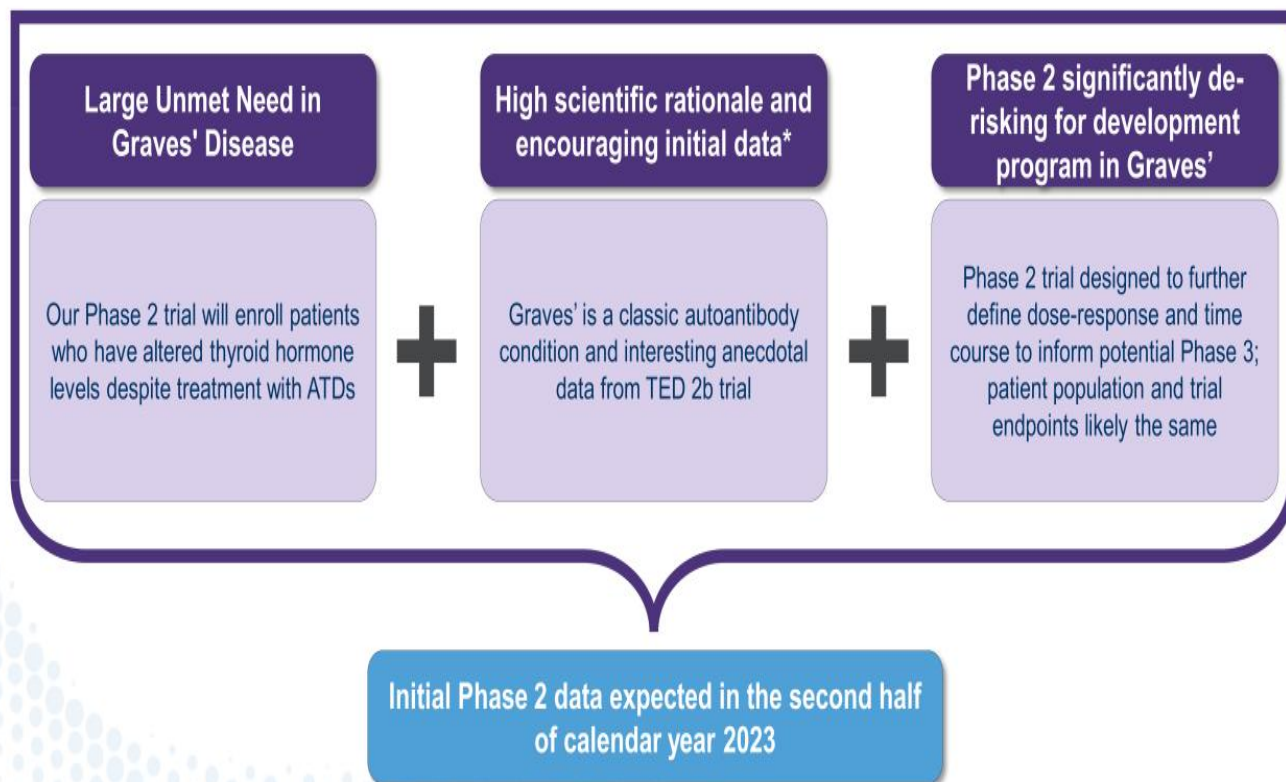
03

FcRn inhibition lowers total and pathogenic IgG and therefore has the potential to significantly improve signs and symptoms in patients who insufficiently respond to anti-thyroid drugs and want to avoid ablative therapy

# Graves' Disease Phase 2 trial measures clinically relevant biomarkers / hormone levels to assess efficacy and to inform a Phase 3 development strategy



## Batoclimab represents a potential targeted therapy for Graves' Disease where unmet need is high



# Closing

# Pipeline expansion: Chronic Inflammatory Demyelinating Polyneuropathy (CIDP) & Graves' Disease

01

CIDP is an exciting opportunity for the anti-FcRn class. We believe that our **uniquely optimized trial** could position batoclimab with best-in-class efficacy and first-to-market simple SC

02

Graves' Disease is a first-in-class opportunity in an autoantibody driven condition with a **straightforward Phase 2** approach and a **high unmet need** in a large subset of patients insufficiently responding even at maximally tolerated doses of current standard of care

03

Data catalysts for new indications complement pivotal data from MG and TED programs. Expect **cadence of data every half year** starting in the second half of 2023

04

Immunovant has \$427M in cash with cash runway into 2025<sup>1,2</sup>



SC = Subcutaneous administration; MG = Myasthenia Gravis; TED = Thyroid Eye Disease

1. As of June 30, 2022, per most recent Quarterly Report on Form 10-Q filed with the SEC on August 5, 2022
2. The assumptions upon which we have based our estimates, including expenditures relating to planned or potential clinical trials, are routinely evaluated and may be subject to change

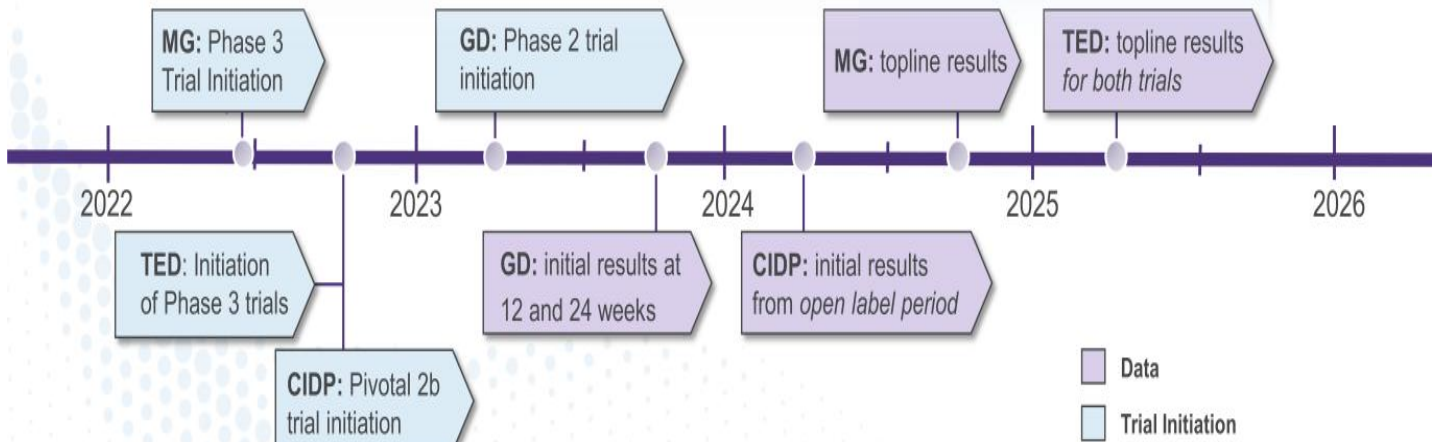
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# Pursuing a broad development program with batoclimab

## Planning for regular cadence of data across indications

Target Indication	Stage of Development
Myasthenia Gravis (MG)	Pivotal Phase 3
Thyroid Eye Disease (TED)	Pivotal Phase 3
Chronic Inflammatory Demyelinating Polyneuropathy (CIDP)	Pivotal Phase 2b*
Graves' Disease (GD)	Phase 2
Warm Autoimmune Hemolytic Anemia (WAIHA)	Phase 2**



\*Registrational package for CIDP may include 1 or 2 pivotal trials depending on a variety of factors

\*\*WAIHA design to be finalized after FDA interaction expected in 2H of calendar 2022

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# Investor Q&A

Pete Salzmann, MD  
Chief Executive Officer

Bill Macias, MD  
Chief Medical Officer





**Immunovant Announces Two New Development Programs for Batoclimab**

- Plan to initiate a Pivotal Phase 2b trial in Chronic Inflammatory Demyelinating Polyneuropathy (CIDP) in the second half of 2022 with initial results from open-label period 1 expected in the first half of 2024
- Plan to initiate a Phase 2 trial in Graves' Disease in early 2023 with initial results expected in the second half of 2023
- Immunovant expects a robust cadence of data every half year beginning in the second half of 2023 through the first half of 2025, including multiple clinical readouts in CIDP, Graves' Disease, Myasthenia Gravis (MG), and Thyroid Eye Disease (TED)
- Company management will host an investor webcast today at 8 AM ET that can be accessed [here](#)

**NEW YORK, September 7, 2022 – Immunovant, Inc.**(Nasdaq: IMVT), a clinical-stage biopharmaceutical company focused on enabling normal lives for people with autoimmune diseases, today announced plans to develop batoclimab in Chronic Inflammatory Demyelinating Polyneuropathy and Graves' Disease.

"We are excited to announce the addition of two new target indications for batoclimab, one of which will be a pivotal program, confirming our confidence in the broad development opportunity for batoclimab", said Pete Salzmann, M.D., Chief Executive Officer of Immunovant. "Evidence suggests that both CIDP and Graves' Disease are caused by autoantibodies and that targeting FcRn is a compelling therapeutic strategy. We believe both indications present promising opportunities ripe for innovation."

"CIDP represents a multi-billion-dollar market for IVIG and a compelling opportunity for the anti-FcRn class, as current therapies for this complex disease have meaningful safety, tolerability and logistical limitations. We have designed our pivotal Phase 2b study leveraging learnings from historical and ongoing clinical trials in this disease, with a goal to improve probability of success and effect size, while studying multiple doses for optimal differentiation" added Bill Macias, M.D., Chief Medical Officer at Immunovant.

"With regard to Graves' Disease, current treatments leave a meaningful proportion of patients unable to achieve normal thyroid hormone function and many remain symptomatic even when on current therapies", said Dr Salzmann. "As a classic autoantibody condition, the straightforward biology of Graves' Disease, from pathogenic autoantibody to altered hormones, provides solid scientific rationale for the indication, supported by anecdotal data from our Thyroid Eye Disease Phase 2b trial. By further defining the dose-response for batoclimab in Graves' Disease, we believe our Graves' Phase 2 trial, if successful, can meaningfully inform and de-risk a future Phase 3 trial and help bring a novel therapy to a large patient population that requires additional treatment."

With the addition of these development programs, Immunovant is now pursuing batoclimab's clinical development in five indications, including MG and TED that have previously disclosed data readouts. Immunovant plans to finalize its trial design in Warm Autoimmune Hemolytic Anemia following interactions with regulators later in 2022.

**Investor Webcast**

Immunovant will host an investor webcast today at 8 AM ET. The webcast will feature prepared remarks by company management and external key opinion leaders and will highlight the current treatment landscape for CIDP and Graves' Disease, as well as plans to study batoclimab's potential to address the unmet needs in its target patient populations. A live question-and-answer session with company management will follow the formal presentations.

Featured speakers will include:

- George Kahaly, MD, PhD, Professor of Medicine and Endocrinology / Metabolism at the Johannes Gutenberg University (JGU) Medical Center
- Jonathan Katz, MD, Director, Neuromuscular Clinic, California Pacific Medical Center
- Todd Levine, MD, Medical Director, Neurology Department, Honor Health Scottsdale, Arizona

To access the webcast, please register [here](#). An archived recording of the webcast will be available on Immunovant's website for a limited time following its conclusion.

**About Immunovant, Inc.**

Immunovant, Inc. is a clinical-stage biopharmaceutical company dedicated to enabling normal lives for people with autoimmune diseases. As a leader in FcRn inhibitor technology, the Company is boldly developing innovative therapies for a range of debilitating autoimmune diseases with significant unmet patient needs. The Company's investigational compound, batoclimab, is a novel, fully human, monoclonal antibody targeting the neonatal Fc receptor (FcRn). For additional information on the Company, please visit [www.immunovant.com](http://www.immunovant.com).

**Forward Looking Statement**

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 "can," "may," "might," "will," "would," "should," "expect," "believe," "estimate," "design," "plan," "intend," and other similar expressions are intended to identify forward-looking statements. Such forward looking statements include Immunovant's plan to initiate a Phase 2b clinical trial for batoclimab in CIDP in the second half of calendar year 2022 with initial results from open-label period 1 expected in the first half of calendar year 2024; Immunovant's plan to initiate a Phase 2 clinical trial for batoclimab in Graves' Disease in early calendar year 2023 with initial results expected in the second half of calendar year 2023; Immunovant's plan to finalize its trial design in Warm Autoimmune Hemolytic Anemia following expected interactions with regulators later in calendar year 2022; Immunovant's plan to develop batoclimab across a broad range of other autoimmune indications; the timing of discussions with regulatory agencies; the size and growth of the potential markets for Immunovant's product candidate and indication selections; Immunovant's expectations regarding timing, the design and results of clinical trials of its product candidates and indication selections; and the potential benefits of batoclimab's unique product attributes. 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; 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

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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, geopolitical tensions, and adverse macroeconomic conditions 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 filed with the SEC on August 5, 2022, 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.

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