
**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**

**January 9, 2023
Date of Report (Date of earliest event reported)**

CABALETTA BIO, INC.
(Exact name of Registrant as Specified in its Charter)

Delaware
(State or other jurisdiction
of incorporation)

001-39103
(Commission
File Number)

82-1685768
(I.R.S. Employer
Identification No.)

**2929 Arch Street, Suite 600,
Philadelphia, PA**
(Address of principal executive offices)

19104
(Zip Code)

(267) 759-3100
(Registrant's telephone number, including area code)

Not Applicable
(Former name or former address, if changed since last report)

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, par value \$0.00001 per share	CABA	The Nasdaq Global Select Market

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.

Item 7.01 Regulation FD Disclosure.

On January 9, 2023, Cabaletta Bio, Inc. (the “Company”) posted to the “Investors & Media” section of the Company’s website at www.cabalettabio.com an updated corporate presentation providing a corporate overview and updated development plan (the “Corporate Presentation”). A copy of the Corporate Presentation is attached hereto as Exhibit 99.1 and is incorporated by reference into this Item 7.01 of this Current Report on Form 8-K.

The information contained in Item 7.01 of this Current Report on Form 8-K, including Exhibit 99.1 attached hereto, is 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 and shall not be incorporated by reference in any filing under the Securities Act of 1933, as amended, or the Exchange Act, except as shall be expressly set forth by specific reference in such filing.

Item 9.01 Financial Statements and Exhibits.**(d) Exhibits**

- 99.1 [Cabaletta Bio, Inc. Corporate Presentation, dated January 9, 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, as amended, the registrant has duly caused this report to be signed on its behalf by the undersigned, hereunto duly authorized.

CABALETTA BIO, INC.

Date: January 9, 2023

By: /s/ Steven Nichtberger
Steven Nichtberger, M.D.
President and Chief Executive Officer

Cabaletta Bio[®]

Corporate Presentation

JANUARY 2023

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Disclaimer

The following presentation, including any printed or electronic copy of these slides, the talks given by the presenters, the information communicated during any delivery of the presentation and any question and answer session and any document or material distributed at or in connection with the presentation (collectively, the "Presentation") has been prepared by Cabaletta Bio, Inc. ("we," "us," "our," "Cabaletta" or the "Company") and is made for informational purposes only. This Presentation does not purport to be a prospectus, to be complete or to contain all of the information you may desire. Statements contained herein are made as of the date of this Presentation unless stated otherwise, and this Presentation shall not under any circumstances create an implication that the information contained herein is correct as of any time after such date or that information will be updated or revised to reflect information that subsequently becomes available or changes occurring after the date hereof. This Presentation may contain "forward-looking statements" within the meaning of the Private Securities Litigation Reform Act of 1995 relating to our business, operations, and financial conditions, and include, but are not limited to, express or implied statements regarding our current beliefs, expectations and assumptions regarding: our business, future plans and strategies for our CAART T and CARTA technologies and CABA™ platform; Cabaletta's ability to grow its autoimmune-focused pipeline; the ability to capitalize on and potential benefits resulting from the translational research partnership with Professor Georg Schett and the exclusive license agreement with IASO Bio; the timing of our planned submission of an Investigational New Drug application (IND) for CABA-201 to the U.S. Food and Drug Administration as well as other planned regulatory filings for our development programs; the progress and results of our DesCAARTes™ Phase 1 trial, including the significance and impact around reported safety and clinical and translational data of cohorts from our DesCAARTes™ trial; the therapeutic potential and clinical benefits of our product candidates; the expectation that Cabaletta may improve outcomes for patients suffering from mucosal pemphigus vulgaris, myasthenia gravis, or other autoimmune diseases; our ability to escalate dosing as high as 10 to 15 billion cells in cohort A6m, initiate dosing in a combination cohort or otherwise; Cabaletta's plans to implement a pre-treatment regimen and the potential ability to enhance *in vivo* DSG3-CAART exposure; our ability to advance dose escalation in the DesCAARTes™ Phase 1 trial at the current dose ranges for the current cohorts and any projected potential dose ranges for future cohorts, and to optimize our targeted cell therapy; our ability to evaluate, and the potential significance of, the relationship between DSG3-CAART persistence and potential clinical responses in patients with mPV; our ability to safely retreat additional patients and whether we will continue to observe a lack of immune-mediated clearance of DSG3-CAART cells after retreatment and repeat dosing of patients; our ability to successfully complete our preclinical and clinical studies for our product candidates, including CABA-201, our ongoing Phase 1 DesCAARTes™ trial, and our ongoing Phase 1 MusCAARTes™ trial of MuSK-CAART, including our ability to enroll the requisite number of patients, dose each dosing cohort in the intended manner, and progress the trial; the ability of MuSK-CAART to target B cells that differentiate into antibody secreting cells, which produce autoantibodies against muscle-specific kinase; our ability to obtain and maintain regulatory approval of our product candidates, including our expectations regarding the intended incentives conferred by and ability to retain Orphan Drug Designation and Fast Track Designation for DSG3-CAART for the treatment of pemphigus vulgaris and Orphan Drug Designation and Fast Track Designation for MuSK-CAART to improve activities of daily living and muscle strength in patients with MuSK antibody-positive myasthenia gravis; the further expansion and development of our modular CABA™ platform across a range of autoimmune diseases; our ability to contract with third-party suppliers and manufacturers, implement an enhanced manufacturing process and further develop our internal manufacturing strategy, capabilities and facilities; our potential commercial opportunities, including value and addressable market, for our product candidates; our expectations regarding our use of capital and other financial results; and our ability to fund operations into the first quarter of 2025. Words such as, but not limited to, "look forward to," "believe," "expect," "anticipate," "estimate," "intend," "plan," "would," "should" and "could," and similar expressions or words, identify forward-looking statements.

Various risks, uncertainties and assumptions could cause actual results to differ materially from those anticipated or implied in our forward-looking statements. Such risks and uncertainties include, but are not limited to, risks related to the success, cost, and timing of our product candidate development activities and preclinical studies and clinical trials, risks related to our ability to demonstrate sufficient evidence of safety, efficacy and tolerability in our preclinical studies and clinical trials of CABA-201, DSG3-CAART and MuSK-CAART, the risk that the results observed with the similarly-designed construct employed in the recent *Nature Medicine* publication are not indicative of the results we seek to achieve with CABA-201, our plans to evaluate additional cohorts in the DesCAARTes™ trial, including a cohort implementing a pre-treatment regimen, the risk that signs of biologic activity or persistence may not inform long-term results, the risk that persistence observed with effective CART-19 oncology studies in combination with lymphodepletion is not indicative of, or applicable to, clinical responses in patients with mPV, risks related to clinical trial site activation or enrollment rates that are lower than expected, our ability to protect and maintain our intellectual property position, risks related to our relationship with third parties, uncertainties related to regulatory agencies' evaluation of regulatory filings and other information related to our product candidates, our ability to retain and recognize the intended incentives conferred by any Orphan Drug Designation and Fast Track Designations, the risk that any one or more of our product candidates will not be successfully developed and commercialized, the risk that the results of preclinical studies or clinical studies will not be predictive of future results in connection with future studies, the impact of COVID-19 on the timing, progress, interpretability of data, and results of ongoing or planned clinical trials and risks relating to as a result of extraordinary events or circumstances such as the COVID-19 pandemic, and any business interruptions to our operations or to those of our clinical sites, manufacturers, suppliers, or other vendors resulting from the COVID-19 pandemic or similar public health crisis. New risks and uncertainties may emerge from time to time, and it is not possible to predict all risks and uncertainties. Except as required by applicable law, we do not plan to publicly update or revise any forward-looking statements contained herein, whether as a result of any new information, future events, changed circumstances or otherwise. Although we believe the expectations reflected in such forward-looking statements are reasonable, we can give no assurance that such expectations will prove to be correct. Accordingly, you are cautioned not to place undue reliance on these forward-looking statements. No representations or warranties (expressed or implied) are made about the accuracy of any such forward-looking statements. For a discussion of these and other risks and uncertainties, and other important factors, any of which could cause our actual results to differ materially from those contained in the forward-looking statements, see the section entitled "Risk Factors" in our most recent annual report on Form 10-K, as well as discussions of potential risks, uncertainties, and other important factors in our other filings with the Securities and Exchange Commission. Certain information contained in this Presentation relates to or is based on studies, publications, surveys and other data obtained from third-party sources and the Company's own internal estimates and research. While the Company believes these third-party sources to be reliable as of the date of this Presentation, it has not independently verified, and makes no representation as to the adequacy, fairness, accuracy or completeness of, any information obtained from third-party sources. The Company is the owner of various trademarks, trade names and service marks. Certain other trademarks, trade names and service marks appearing in this Presentation are the property of third parties. Solely for convenience, the trademarks and trade names in this Presentation are referred to without the ® and TM symbols, but such references should not be construed as any indicator that their respective owners will not assert, to the fullest extent under applicable law, their rights thereto.

Develop and launch the first curative
targeted cellular therapies for patients
with autoimmune diseases

Cabaletta Bio[®]

Cabaletta®: Pursuing cures for a broad range of autoimmune diseases

Recent expansion into CD19-CAR T complements & leverages deep cell therapy experience in autoimmunity

CARTA Strategy | CABA-201 (4-1BB CD19-CAR T) on track for 1H23 IND submission

- **Builds on academic clinical data¹ revealing potential for CD19-CAR T to reset the immune system in refractory SLE patients**
 - *Exclusive translational research partnership with lead investigator¹ providing early & actionable insights for CABA-201*
- **CABA-201 has been specifically engineered for patients with autoimmune diseases**
 - *Including the same 4-1BB costimulatory domain¹ and similar CD19 binder affinity² as used in the academic SLE study*
 - *Fully human CD19 binder in CABA-201 with a favorable clinical tolerability profile in ~20 oncology patients*
- **Potential to cure many autoimmune diseases such as SLE, RA, myositis and systemic sclerosis³**

CAART Strategy | DSG3-CAART & MuSK-CAART clinical studies evaluating combination regimens

- **DesCAARTes™ trial in mucosal pemphigus vulgaris – 1 month safety & persistence data anticipated 1H23⁴**
 - *Enrolling in combination sub-study using pre-treatment with IVIg & cyclophosphamide*
- **MusCAARTes™ trial in MuSK myasthenia gravis – leveraging insights from autoimmune experience with DSG3-CAART**
 - *Initiated in 4Q22; received FDA Fast Track Designation & Orphan Drug Designation*

Initial CABA-201 clinical data⁵ and 6-month combination data from CAART trials expected by 1H24⁴

CAART – Chimeric AutoAntibody Receptor T cells; CARTA – Chimeric Antigen Receptor T cells for Autoimmunity; IND – Investigational New Drug; SLE – Systemic lupus erythematosus; RA – Rheumatoid arthritis

1. Mackensen, Andreas, et al. "Anti-CD19 CAR T cell therapy for refractory systemic lupus erythematosus." *Nature Medicine* (2022): 1-9.

2. Dai, Zhenyu, et al. "Development and functional characterization of novel fully human anti-CD19 CARs for T-cell therapy." *Journal of Cellular Physiology* 236.8 (2021): 5832-5847.

3. Mullin, Emily. "How a 'Living Drug' Could Treat Autoimmune Disease." *WIRED*, 16 Sept 2022.

4. Assumes no dose-limiting toxicities are observed in the cohort, uninterrupted enrollment and no delays due to COVID-19 resurgence occur in the trial.

5. Subject to and pending clearance of CABA-201 IND by the FDA.

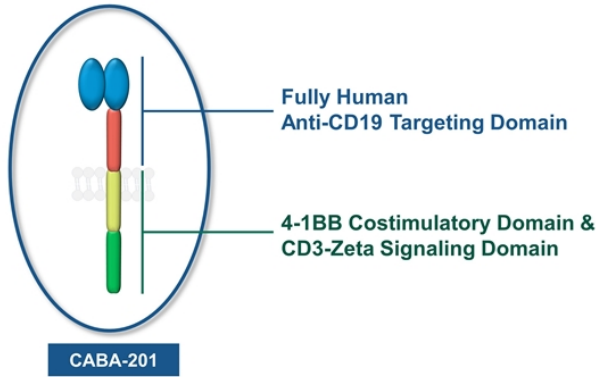
One CABA™ platform, two strategies to address autoimmune diseases

Complementary strategies to optimize clinical outcomes using cellular therapies in autoimmune diseases

CARTA

Chimeric Antigen Receptor T cells for Autoimmunity

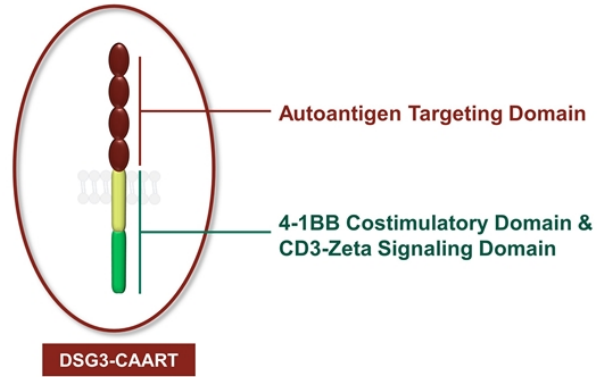
Potential to **'reset the immune system'** in patients with **autoimmune diseases driven by B cells**, through generalized transient B cell depletion and **repopulation of healthy B cells**¹



CAART

Chimeric AutoAntibody Receptor T cells

In autoimmune diseases with a **limited number of well-defined pathogenic autoantibodies**, permanent **antigen-specific B cell depletion** may provide an **elegant biologic solution** to disease²



1. Mackensen, Andreas, et al. "Anti-CD19 CAR T cell therapy for refractory systemic lupus erythematosus." *Nature Medicine* (2022): 1-9.
2. Ellebrecht, Christoph T., et al. "Reengineering chimeric antigen receptor T cells for targeted therapy of autoimmune disease." *Science* 353.6295 (2016): 179-184.

CABA™ platform may apply across a range of autoimmune diseases

Biologic opportunity for cure or treatment may be possible in dozens of B-cell mediated autoimmune diseases*

- **Myasthenia Gravis**^{1,2,3,5}
- Multiple Sclerosis⁶
- Neuromyelitis Optica³
- Chronic Inflammatory Demyelinating Polyneuropathy^{1,2}
- Anti-NMDAR Encephalitis^{3,6}
- Lambert-Eaton Syndrome⁵

Neurology

Rheumatology

- **Systemic Lupus Erythematosus**^{3,4,5,6}
- Rheumatoid Arthritis^{2,3,4}
- **Myositis**⁵
- **Systemic sclerosis**⁶
- ANCA-Associated Vasculitis^{3,4,5}

- **Pemphigus Vulgaris**^{1,2,3}
- Pemphigus Foliaceus^{1,2,3}
- Epidermolysis Bullosa Acquisita³
- Bullous Pemphigoid^{1,2,3}

Dermatology

Hematology

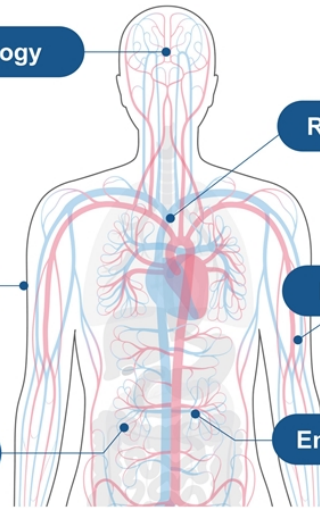
- Immune Thrombocytopenic Purpura³
- Thrombotic Thrombocytopenic Purpura^{1,2,3}
- Antiphospholipid Syndrome^{4,5}
- Autoimmune Hemolytic Anemia³

- **Lupus Nephritis**^{3,4}
- Membranous Nephropathy^{1,2,3}
- Goodpasture's Syndrome^{1,2,3,4}

Nephrology

Endocrinology

- Type 1 Diabetes^{3,6}
- Graves' Disease^{3,5}
- Hashimoto's Disease⁵



* Illustrative list of diseases where biologic opportunity for cure or treatment with CAART or CARTA approach may be possible.

Diseases **in bold** represent where clinical studies are underway or plan to be initiated by Cabaletta or by Professor Schett

1. Konecny, Inga. "Update on IgG4-mediated autoimmune diseases: new insights and new family members." *Autoimmunity Reviews* (2020): 102646.

2. Huijbers, Maartje G., et al. "IgG4-mediated autoimmune diseases: a niche of antibody-mediated disorders." *Annals of the New York Academy of Sciences* 1413.1 (2018): 92.

3. Ludwig, Ralf J., et al. "Mechanisms of autoantibody-induced pathology." *Frontiers in immunology* 8 (2017): 603.

4. Suurmond, Jolien, and Betty Diamond. "Autoantibodies in systemic autoimmune diseases: specificity and pathogenicity." *The Journal of clinical investigation* 125.6 (2015): 2194-2202.

5. Xiao, Ze Xiu, Joseph S. Miller, and Song Guo Zheng. "An updated advance of autoantibodies in autoimmune diseases." *Autoimmunity Reviews* (2020): 102743.

6. Hampe, Christiane S. "B cells in autoimmune diseases." *Scientifica* 2012 (2012).

Pipeline targeting autoimmune diseases where cure is possible

CABA™ Platform	Indication	Program	Discovery	Preclinical	Phase 1	Phase 2/3
CARTA <i>Chimeric Antigen Receptor T cells for Autoimmunity</i>	Multiple Undisclosed Indications	CABA-201 <i>4-1BB CD19-CAR T</i>				
CAART¹ <i>Chimeric AutoAntibody Receptor T cells</i>	Mucosal Pemphigus Vulgaris	DSG3-CAART				
	MuSK Myasthenia Gravis	MuSK-CAART				
	PLA2R Membranous Nephropathy	PLA2R-CAART				
	Mucocutaneous Pemphigus Vulgaris	DSG3/1-CAART				

1. Additional disease targets in discovery stage in our CAART pipeline portfolio include hemophilia and two undisclosed indications.



Chimeric Antigen Receptor T Cells for Autoimmunity
CABA-201

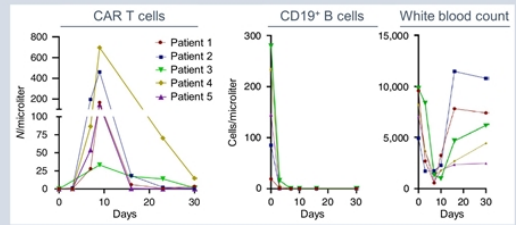
Cabaletta Bio®

Academic clinical data: Immune system reset in 5/5 SLE patients¹

Exclusive translational research partnership with lead investigator informing our CD19 clinical strategy

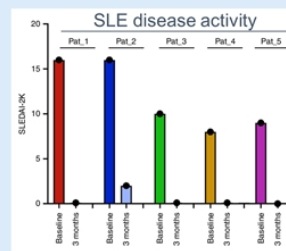
5/5 refractory SLE patients treated with 4-1BB CD19-CAR T² resulting in rapid, deep and transient depletion of CD19⁺ B cells¹

- All 5 patients with moderate to severe disease
- Preconditioning with standard Flu/Cy regimen³
- Dose of 1×10^6 CD19-CAR T cells/kg
- CD19 binder²: 4-1BB costimulatory domain & FMC63 scFv



Clinical & serologic responses by 3 mo. after 4-1BB CD19-CAR T therapy with promising safety profile¹

- Anti-dsDNA antibodies undetectable in 5/5
 - All SLE-associated antibodies reduced
 - Complement levels normalized
- No – or only mild – CRS observed
 - Grade 1 fever in 3/5 patients
- No neurotoxicity / ICANS



Repopulation of healthy B cells¹

- New, immature B cells reappeared in 5/5 patients between 2-5 months
- Limited decline in vaccination titers

Durable clinical responses¹

- 5-17 months of follow up
- Responses maintained with no need for SLE-associated medications

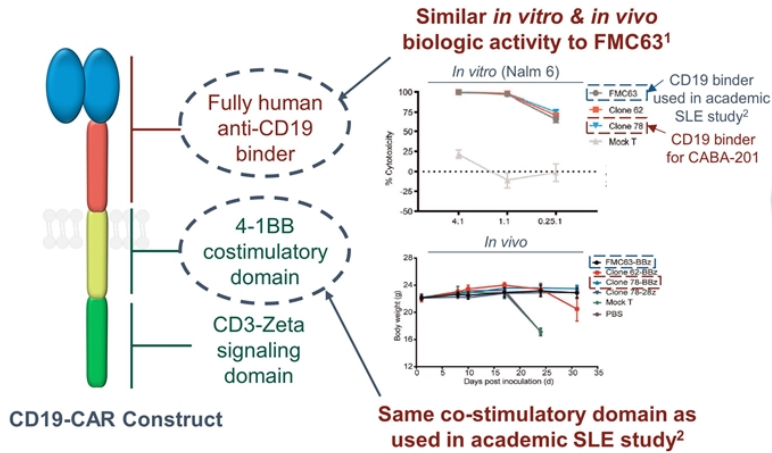
SLE – Systemic lupus erythematosus; scFv – Single chain variable fragment; SLEDAI-2K – Systemic Lupus Erythematosus Disease Activity Index 2000; CRS – Cytokine release syndrome; ICANS – Immune effector cell-associated neurotoxicity syndrome

1. Mackensen, Andreas, et al. "Anti-CD19 CAR T cell therapy for refractory systemic lupus erythematosus." *Nature Medicine* (2022): 1-9.
2. The construct utilized in this study has a similar design to CABA-201, sharing the 4-1BB costimulatory domain, but is a different construct.
3. Fludarabine (Flu) 25 mg/m²/d intravenously day -5 to day -3; Cyclophosphamide (Cy) 1,000 mg/m²/d intravenously on day -3

CABA-201: CD19-targeting CAR T therapy for autoimmune diseases

Cabaletta's CD19 binder with similar *in vitro* & *in vivo* activity to FMC63¹ (binder used in academic SLE study²)

CABA-201



Clinical Data for Licensed CD19 Binder³

Fully human binder

Evaluated within dual-CAR combined with CD22 binder with standard Flu/Cy preconditioning

Evaluated in ~20 patients to date³

Under evaluation in patients with B cell leukemia and lymphoma in IIT in China

Promising tolerability data to date³

Used in oncology patients with high target cell burden

SLE – Systemic lupus erythematosus; IIT – Investigator-initiated trial; Flu/Cy – Fludarabine/Cyclophosphamide; CRS – Cytokine release syndrome; ICANS – Immune effector cell-associated neurotoxicity syndrome

1. Dai, Zhenyu, et al. "Development and functional characterization of novel fully human anti-CD19 chimeric antigen receptors for T-cell therapy." *Journal of Cellular Physiology* 236.8 (2021): 5832-5847.

2. Mackensen, Andreas, et al. "Anti-CD19 CAR T cell therapy for refractory systemic lupus erythematosus." *Nature Medicine* (2022): 1-9.

3. Evaluated as part of CT120, a dual-CD19xCD22 CAR T product candidate under development by Nanjing IASO Biotherapeutics, Co., Ltd. (IASO Bio).

Accelerating development of CABA-201 for autoimmune diseases

Differentiated asset, exclusive translational insights & track record of timely CAART clinical implementation

1 Cabaletta is uniquely positioned to efficiently advance CABA-201 in autoimmune diseases

- Expertise in conducting complex, interdisciplinary autoimmune-focused cell therapy clinical trials
- Track record of positive regulatory interactions to support cell trials in autoimmune diseases since 2018
 - 2 INDs cleared within routine 30-day period; multiple clinical protocol modifications, Fast Track designations granted
- Successful track record manufacturing novel cell therapy products with academic and industry partners
- Deep understanding of autoimmunity allows potential application across broad range of diseases


2 Clinically-evaluated binder & translational research partnership support CABA-201 development

- CABA-201 leverages similar design to 4-1BB-containing CD19-CAR T in seminal academic SLE study¹
- Exclusive research partnership with lead investigator for SLE study provides early and actionable insights

IND submission for CABA-201 anticipated in first half of 2023

SLE – Systemic lupus erythematosus

1. Mackensen, Andreas, et al. "Anti-CD19 CAR T cell therapy for refractory systemic lupus erythematosus." *Nature Medicine* (2022): 1-9.

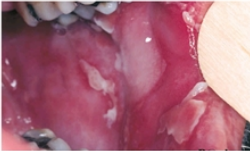



Chimeric AutoAntibody Receptor T Cells
DSG3-CAART & MuSK-CAART

Cabaletta Bio®

Overview of pemphigus vulgaris & current treatment landscape

Current treatments require broad immunosuppression associated with safety risks and transient efficacy

	Mucosal PV ¹ (25%)	Mucocutaneous PV ² (75%)
		
Associated Antibody	Anti-DSG3	Anti-DSG3 + Anti-DSG1
Clinical Signs	Painful blisters of the mucous membranes (mouth, nose, larynx, esophagus, eyes, genitalia, rectum)	Blisters on orifices and skin
U.S. Disease Prevalence	3,250 to 4,750	9,750 to 14,250

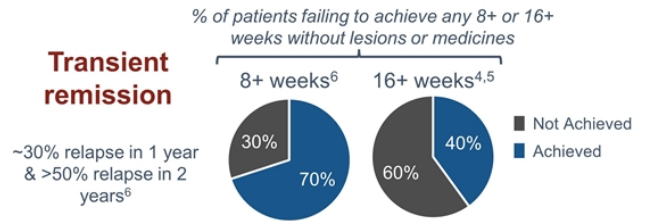
CROT = 8+ weeks without lesions while off systemic therapy

1. Image credit: D@nderm.
 2. <http://www.vgrd.org/archive/cases/2004/pv/DSCN4996%20copy.JPG>
 3. Joly, Pascal, et al. "First-line rituximab combined with short-term prednisone versus prednisone alone for the treatment of pemphigus (Ritux 3): a prospective, multicentre, parallel-group, open-label randomised trial." *The Lancet* 389.10083 (2017): 2031-2040.
 4. Werth, Victoria P., et al. "Rituximab versus Mycophenolate Mofetil in Patients with Pemphigus Vulgaris." *New England Journal of Medicine* (2021).
 5. Rituximab label, 08/2020 revision.
 6. Kushner, Carolyn J., et al. "Factors Associated With Complete Remission After Rituximab Therapy for Pemphigus." *JAMA dermatology* (2019).
 7. Tony, Hans-Peter, et al. "Safety and clinical outcomes of rituximab therapy in patients with different autoimmune diseases: experience from a national registry (GRAID)." *Arthritis research & therapy* 13.3 (2011): 1-14.

Broad immunosuppression^{3,6}

Modestly effective & poorly tolerated

Rituximab plus steroids (~3,500 mg/yr)⁴



- Safety risks**
- 22% annual serious adverse event (SAE) rate⁴
 - 4-9%^{3,4,5} annual risk of severe infection in PV
 - ~1.9% lifetime risk of fatal infection⁷

Ongoing DesCAARTes™ study in patients with mucosal PV

Monotherapy DSG3-CAART demonstrates favorable tolerability to date, but persistence plateaus



DesCAARTes™ study of DSG3-CAART

Ongoing open-label Phase 1 study¹ to determine the maximum tolerated dose & to evaluate safety of DSG3-CAART

Part A Cohorts ¹	Subjects	Dose*
A1 – A6m ^{2,3}	3 (+3) per cohort	20M to 15B

Primary objective:

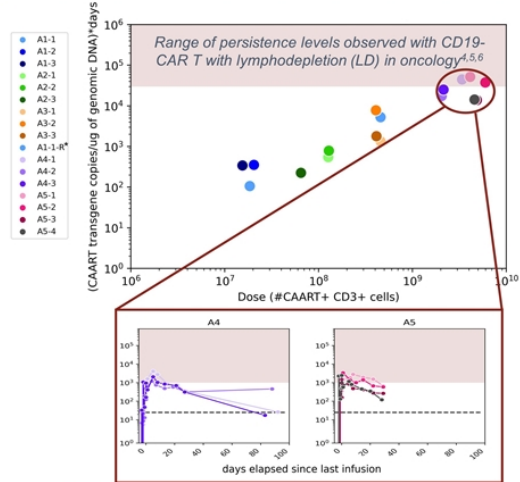
Determine the maximum tolerated dose of DSG3-CAART

Primary endpoint:

Adverse events, including dose-limiting toxicities (DLTs), related to DSG3-CAART within 3 months of infusion

Combination Sub-study	Subjects	Dose*
IVIg / Cyclophosphamide ³	3 (+3) per cohort	2.5B

DSG3-CAART Persistence to 29d in Cohorts A1-A5



* A1-1-R refers to a subject in cohort A1 who was retreated at the cohort A3 dose (500M).

1. FDA has requested, and the Company has agreed, that we will share data from part A to inform a discussion on the optimal design of part C. According to FDA advice, the submission of part A data is not gating to planned enrollment in part B.
 2. Cohort A6m reflects a two-dose regimen of 5.0 to 7.5 billion cells, 3 weeks apart, for a total of 10 to 15 billion cells, which is designed to prolong exposure time of DSG3-CAART in vivo.
 3. Combination cohort has been prioritized relative to A6m based on emerging data in cohorts A4 and A5.
 4. Mueller, Karen Thudium, et al. "Cellular kinetics of CTL019 in relapsed/refractory B-cell acute lymphoblastic leukemia and chronic lymphocytic leukemia." *Blood*, 130.21 (2017): 2317-2325.
 5. Schuster SJ, Bishop MR, Tam CS, et al. Tisagenlecleucel in Adult Relapsed or Refractory Diffuse Large B-Cell Lymphoma. *N Engl J Med*. 2019;380(1):45-56. doi:10.1056/NEJMoa1804980.
 6. The range of persistence observed with anti-CD19 CART therapy in oncology has not been confirmed to be necessary or sufficient for clinical responses in patients with mPV.
 * 20M, 100M, 500M, 2.5B, 5.0B to 7.5B and 10B to 15B refers to the number of DSG3-CAART cells infused for patients in dosing cohorts A1 to A6m (M – millions; B – billions).

Combination sub-study prioritized to increase CAART exposure

Combination strategy designed to enhance cytokine & diminish autoantibody effects on CAART activity

Combination sub-study cohort

A4 dose (2.5×10^9 cells) + cyclophosphamide (CY) & IVIg

- Dose-dependent increase in CAART persistence as monotherapy plateaued with Cohort A5
 - Through up to 6 months post-CAART infusion, no clear pattern in antibody levels and disease activity observed in first 3 subjects at cohort A5 dose
- CY may reduce 'cytokine sink,' potentially enhancing CAART activation & proliferation
- CY + IVIg may reduce anti-DSG3 autoantibodies, addressing a potential efficacy barrier
- CY & IVIg likely to provide transient improvement in first few months after infusion^{1,2,3,4,5}
 - DSG3-CAART clinical effect may require follow-up for 6-9 months

Cohort A6m | *2x A5 dose ($1-1.5 \times 10^{10}$ cells) – lower priority*

- Two infusions at the A5 dose level 3 weeks apart
 - To potentially increase the duration of maximal exposure to DSG3-CAART

1. Amagai, Masayuki, et al. "A randomized double-blind trial of intravenous immunoglobulin for pemphigus." *Journal of the American Academy of Dermatology* 60.4 (2009): 595-603.
 2. Arnold, D. F., et al. "An 'n-of-1' placebo-controlled crossover trial of IVIg as adjuvant therapy in refractory pemphigus vulgaris." *British Journal of Dermatology* 160.5 (2009): 1098-1102.
 3. Zhang, Wenjing, et al. "Short-Term Intravenous Infusion of Cyclophosphamide in the Treatment of Refractory Pemphigus Vulgaris: A Retrospective Study." *Dermatology* 237.2 (2021): 185-190.
 4. Fleischli, Mary E., Rachel H. Valek, and Amit G. Pandya. "Pulse intravenous cyclophosphamide therapy in pemphigus." *Archives of dermatology* 135.1 (1999): 57-61.
 5. Lois, Margarita, et al. "Effect of IVIg with or without cytotoxic drugs on pemphigus intercellular antibodies." *Journal of the American Academy of Dermatology* 64.3 (2011): 484-489.

High unmet need in MuSK myasthenia gravis

Strategy for first-in-human trial informed by learnings from DesCAARTes™ study

Orphan Drug Designation

Fast Track Designation

1 Compelling biologic rationale, similar to PV

- IgG4-dominant disease
- Autoantibody titers drop after rituximab^{1,2}
- Pathogenic B cells incompletely eliminated by rituximab and persist during relapse³
- Circulating anti-MuSK antibody titers may be ~90% lower than anti-DSG3 antibody in PV^{4,5,6}

2 Differentiated market opportunity

- Total U.S. MG prevalence 50,000 to 80,000 patients
- MuSK+ MG comprises 6-7.5% of total MG population
- Compared to AChR+ MG, MuSK+ disease is typically more severe with limited treatment options
- MuSK+ disease has early onset, 7:1 females

* 500M refers to the number of MuSK-CAART cells infused for patients in dosing cohort A1 (M – millions).

1. Hain, Berit, et al. "Successful treatment of MuSK antibody-positive MG with rituximab." *Muscle & Nerve*. 33.4 (2006): 575-580.

2. Ila, Isabel, et al. "Sustained response to Rituximab in anti-AChR and anti-MuSK positive Myasthenia Gravis patients." *Journal of neuroimmunology* 201 (2008): 90-94.

3. Jiang, Ruoyi, et al. "Single-cell repertoire tracing identifies rituximab-resistant B cells during myasthenia gravis relapses." *JCI insight* 5.14 (2020).

4. Matthews, Ian, et al. "Muscle-specific receptor tyrosine kinase autoantibodies—a new immunoprecipitation assay." *Clinica chimica acta* 348.1-2 (2004): 95-99.

5. McConville, John, et al. "Detection and characterization of MuSK antibodies in seronegative myasthenia gravis." *Annals of neurology* 55.4 (2004): 580-584.

6. Marino, Mariapaola, et al. "Long-lasting rituximab-induced reduction of specific—but not total—IgG4 in MuSK-positive myasthenia gravis." *Frontiers in immunology* 11 (2020): 613.

7. A total of 6 subjects will need to have received the final selected dose in Part A of the study.

8. Cohorts A3 and A4 will be enrolled concurrently after cohort A2 is well-tolerated with a preference for enrollment into A4.

MusCAARTes™ study of MuSK-CAART

Ongoing open-label study to determine the maximum tolerated dose & to evaluate safety of MuSK-CAART

Part	Cohort	Subjects
A – Monotherapy Dose Escalation⁷ Dose increasing from 500M with 2 (+4) design	A1-A3	2 (+4) per cohort
A – Adaptive Combination Cohorts⁷ Combination cohorts ⁸ , starting at A2 dose	A4+	2 (+4) per cohort
B – Expansion Expanded subject enrollment at final selected dose	B	~12

Study Endpoint & Objectives

Primary Endpoint: Adverse Events, including DLT

Secondary & Tertiary Objectives: CAAR T expansion/persistence, effect on serum anti-MuSK antibody titer, use of concomitant systemic medications, effect on clinical symptoms, manufacturing success rate



Corporate Summary

Cabaletta Bio[®]

Manufacturing strategy

Three-stage approach allows for efficient allocation of capital while leveraging experienced partners



1. CDMOs shown are currently contracted for select CAART product candidates.

Cabaletta Bio leadership

LEADERSHIP TEAM



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President, CEO & Chairman



Samik Basu, M.D.
Chief Scientific Officer



Gwendolyn Binder, Ph.D.
President, Science & Technology



David J. Chang, M.D., M.P.H., FACR
Chief Medical Officer



Arun Das, M.D.
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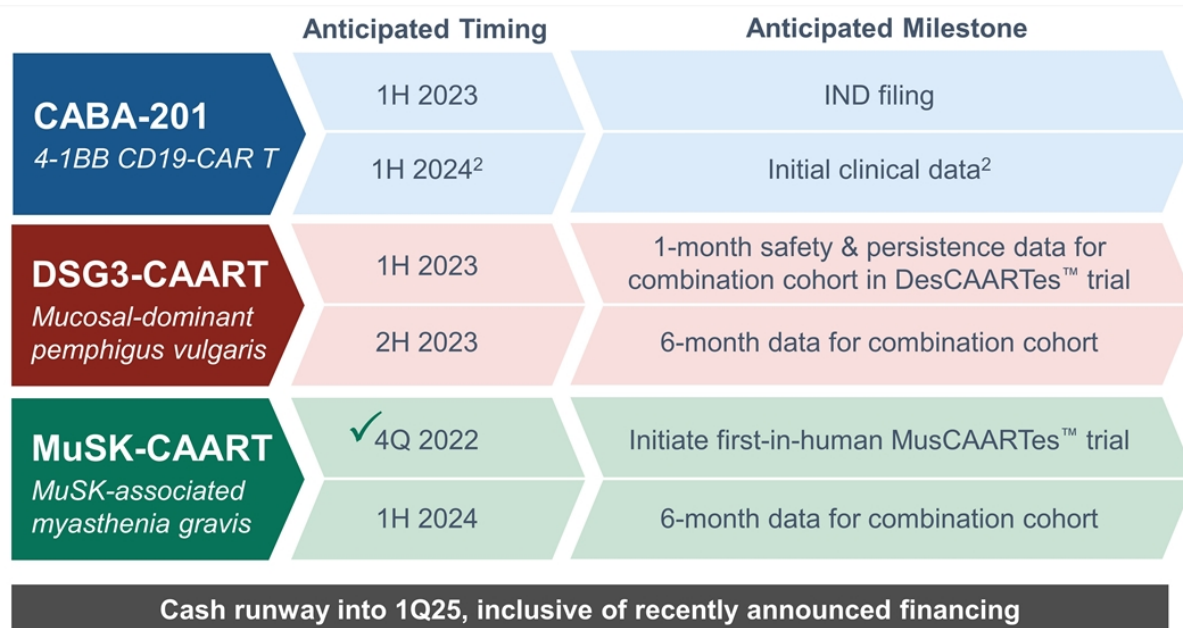
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Track record of operational success employing novel cell therapies in autoimmunity across preclinical, clinical, manufacturing & regulatory domains

Multiple potential clinical catalysts anticipated in next 12-18 months¹



1. Assumes no dose-limiting toxicities are observed in any cohort, uninterrupted enrollment and no delays due to COVID-19 resurgence occur in the trial.
 2. Subject to and pending clearance of CABA-201 IND by the FDA.

Cabaletta Bio[®]

A microscopic view of several spherical cells with a red, textured interior, set against a white background. The cells are out of focus, with one in the foreground being sharper.

Corporate Presentation

JANUARY 2023

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