UNITED STATES SECURITIES AND EXCHANGE COMMISSION

Washington, DC 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 13, 2025

SAVARA INC.

(Exact name of registrant as specified in its charter)

Delaware (State or other jurisdiction of incorporation) 001-32157 (Commission 84-1318182 (IRS Employer Identification No.)

1717 Langhorne Newtown Road, Suite 300
Langhorne, PA 19047
(Address of principal executive offices, including zip code)

(512) 614-1848

(Registrant's telephone number, including area code)

 $\label{eq:N/A} N/A \end{result}$ (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 (see General Instruction A.2. below):

| Ш | 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 240.14c |

☐ 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:

| | Trading | Name of each exchange |
|---|-----------|---------------------------------|
| Title of each class | Symbol(s) | on which registered |
| Common Stock, par value \$0.001 per share | SVRA | The Nasdaq Global Select Market |

Indicate by check mark whether the registrant is an emerging growth company as defined in as defined in Rule 405 of the Securities Act of 1933 (\S 230.405 of this chapter) or Rule 12b-2 of the Securities Exchange Act of 1934 (\S 240.12b-2 of this chapter).

Emerging growth company \square

If an emerging growth company, indicate by check mark if the registrant has elected not to use the extended transition period for complying with any new or revised financial accounting standards provided pursuant to Section 13(a) of the Exchange Act. \Box

Item 7.01. Regulation FD Disclosure.

Savara has updated its corporate presentation, which is available on the Investor Relations page of Savara's website at https://savarapharma.com/investors/events-presentations/. A copy of the presentation is furnished as Exhibit 99.1 to this Current Report on Form 8-K. Savara undertakes no duty or obligation to update or revise the information contained in this presentation, although it may do so from time to time. Any such updates may be made through the Investor Relations page of the Savara website, the filing of other reports or documents with the U.S. Securities and Exchange Commission (the "SEC"), press releases, or other public disclosure.

The information in Item 7.01 in this Current Report on Form 8-K shall not be deemed "filed" for purposes of Section 18 of the Exchange Act, or otherwise subject to the liabilities of that Section, nor shall it 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.

Exhibit No. Description

99.1 Savara Corporate Presentation

104 Cover Page Interactive Data File (embedded within the Inline XBRL document).

SIGNATURES

Pursuant to the requirements of the Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned hereunto duly authorized.

Date: January 13, 2025

SAVARA INC. a Delaware corporation

By:

/s/ Dave Lowrance
Dave Lowrance
Chief Financial & Administrative Officer



Corporate Overview

Developing New Therapies for Rare Respiratory Diseases January 2025

Safe Harbor Statement

Savara Inc. ("Savara" or the "Company") cautions you that statements in this presentation that are not a description of historical fact are forward-looking statements which may be identified by the use of words such as "expect," "intend," "plan," "anticipate," "believe," and "will," among others. Such statements include, but are not limited to, statements regarding the potential health benefits and risks and projected development timeline of MOLBREEVI; the timing of regulatory submissions; the potential for and impact of regulatory approval; the potential addressable patient population, market size, commercial opportunity, and competitive landscape for MOLBREEVI; Savara's commercial launch planning activities, including disease awareness campaign, GM-CSF autoantibody testing, planned infrastructure, and anticipated hiring and the potential impact of those activities; and the sufficiency of our resources to fund the advancement of our development program and potential sources of additional capital. Savara may not actually achieve any of its plans or product development goals in a timely manner, if at all, or otherwise carry out its current intentions or meet the expectations or projections disclosed in its forward-looking statements, and you should not place undue reliance on these forward-looking statements. These forward-looking statements are based upon Savara's current expectations and involve assumptions that may never materialize or may prove to be incorrect. Actual results and the timing of events could differ materially from those anticipated in such forward-looking statements as a result of various risks and uncertainties, which include, without limitation, the risks associated with our ability to successfully develop, obtain regulatory approval for and commercialize MOLBREEVI for aPAP; the risks and uncertainties related to the impact of widespread health concerns and geopolitical conditions on our business and operations; risks and uncertainties associated with the ability to project future cash utilization and reserves needed for contingent future liabilities and business operations; the ability to successfully conduct clinical trials for our product candidate; the availability of sufficient resources and the timing and ability of Savara to raise additional capital as needed to fund continued operations. The risks and uncertainties facing Savara are described more fully in Savara's filings with the Securities and Exchange Commission including our filings on Form 8-K, our Annual Report on Form 10-K for the fiscal year ended December 31, 2023, and our Quarterly Report on Form 10-Q for the quarter ended Sept. 30, 2024.

You are cautioned not to place undue reliance on our forward-looking statements, which speak only as of the date on which they were made. Savara undertakes no obligation to update such statements to reflect events that occur or circumstances that exist after the date on which they were made, except as may be required by law. Third-party information included herein has been obtained from sources believed to be reliable, but the accuracy or completeness of such information is not guaranteed by, and should not be construed as a representation by, the Company. Additionally, this presentation includes internal research and estimates performed by the Company, which have not been independently verified.

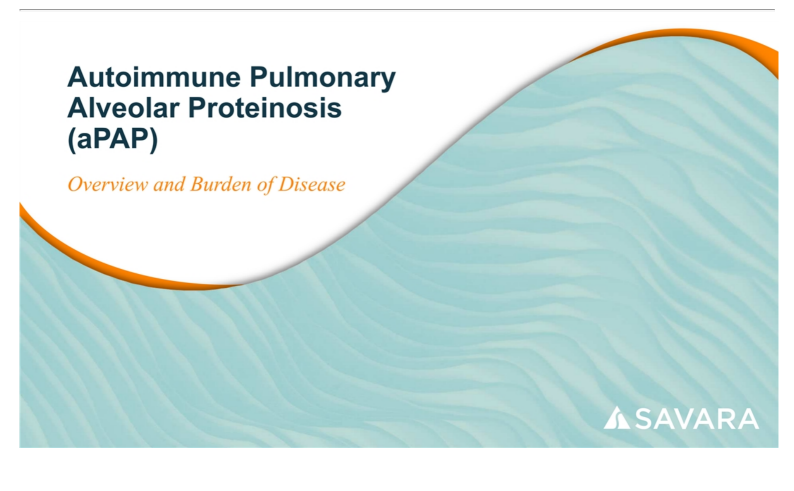
MOLBREEVI (molgramostim inhalation solution) is an investigational product that has not been approved for sale or determined to be safe or effective by the U.S. Food & Drug Administration or any regulatory authority.

MOLBREEVI, MY MOLBREEVI and aPAP ClearPath are trademarks of Savara. All other trademarks included herein are the property of the owners thereof and are used for reference purposes only.

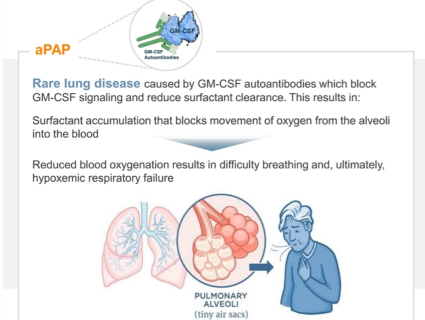
Executive Leadership Team

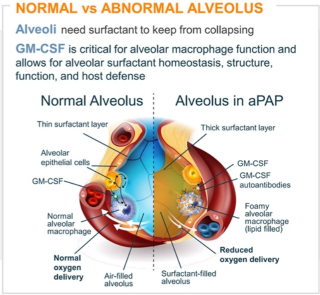


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Autoimmune PAP: Disease of Alveolar Macrophage Dysfunction





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Autoimmune PAP is a Rare, Long-Term, Chronic Lung Disease

No approved drugs in the U.S. or Europe for aPAP, only treatment option is an invasive procedure

Progressive Shortness of Breath

- Gas exchange in the lungs is impaired and patients may experience shortness of breath
- At first it occurs upon exertion, but as disease progresses, it can occur even when a person is at rest



Increased Risk of Infection

 Serious infections, the most common and threatening complications of aPAP, occur in 5–13% of patients and account for 18–20% of deaths¹⁻⁴



Fatigue, Decreased Exercise Tolerance

 Fatigue and significantly reduced exercise capacity can dramatically impact the simplest of daily activities, e.g., getting winded walking up a flight of stairs



Fibrosis and Lung Transplant

 Over time, aPAP can lead to pulmonary fibrosis and respiratory failure which can be fatal and may require lung transplantation



Cough and Episodes of Fever

 Cough, sputum production, and episodes of fever, especially if secondary lung infection develops

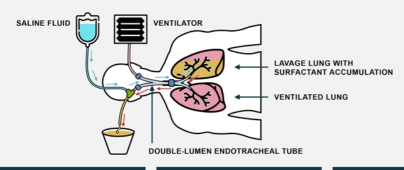


1. Trapnell Nat Rev Dis Primers 2019; 2. Seymour AJRCCM 2002; 3. Inoue AJRCCM 2008; 4. Jouneau Respirology 2020

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Whole Lung Lavage (WLL):

- Performed under anesthesia and requires hospitalization, a team of experienced HCPs, and surgical resources
- Does not correct underlying pathophysiology of the disease or prevent abnormal surfactant accumulation and often needs to be repeated
- Patients describe WLL as burdensome and emotionally taxing
- Long-term negative impact (potential lung damage) of repeated WLL procedures is unknown



Requires insertion of doublelumen endobronchial tube for lung separation

Treated lung is repeatedly filled with up to 15-50L of saline and then drained by gravity

Patient is percussed to emulsify the surfactant sediment

continued until lavage fluid becomes clear

Saline is drained by gravity and

A Lung Lavage is an

Performed in a Tertiary

Invasive Procedure

Center and is Not

Standardized

Sources: 1: Campo, Assessment and Management of PAP in a Reference Center, Orphanet Jour. of Rare Dis., 2013; 2: Campo, Nat. History of PAP Data from Italian Nat. Reference Center, ERJ, 2019; Seymour, J. J. Pulmonary alveolar proteinosis: Progress in the First 44 Years, Am. J. Respir Crit. Care Med, 2002; 3: Udwadia, Jain. NEJM (2007) 357:19, 4 McCarthy, Autoimmune Pulmonary Alveolar Proteinosis, Amer. Journal of Respiratory and Critical Care Med., 2022.

Disease Burden: Autoimmune PAP Patients Have Significantly Higher Rates of Healthcare Utilization and Comorbidities¹



Charlson **Comorbidity Index** (CCI)*



PAP: 1.84 ± 2.48

Age and Gender Matched Controls: 0.55 ± 1.44

P value: <0.0001

*Developed to classify comorbid conditions which may influence mortality risk. Most widely used comorbidity index used to determine survival rates in patients with multiple comorbidities.





Age and Gender Matched Controls: 10.40 ± 11.38

Outpatient visits

(~17 per year)

P value: <0.01





Emergency Room Visits

(~1.5 per year)

+38% Vs.

PAP: 1.49 ± 1.17

Age and Gender Matched Controls: 1.08 ± 0.27

P value: 0.014



Longer hospital stays

(~16 days per year)



PAP: 15.96 ± 20.71

Age and Gender Matched Controls: 5.40 ± 5.07

P value: 0.027

1: McCarthy C, et al. Orphanet Journal of Rare Diseases (2018) 13:129

Patient Perspective on Living with aPAP

66

With whole lung lavage being the only treatment option, it's terrifying. The best way to describe it in layman's terms, it's like a car wash for your lungs. Having an alternative treatment from whole lung lavage would mean the world to me, it would give me the opportunity to get my life back. To give me the freedom of what I had before aPAP.

- Kelsea

"

66

Overall, when the surfactant builds up, I notice how much more tired I get, walking from the basement to the first floor will wind me, I'll get chest congestion and cough up yellow mucus. So, every 8 months surfactant builds up and I'll need the whole lung lavage, and it causes a lot of anxiety knowing I will need to keep having them. Having had multiple lung lavages over the years; there needs to be more options when it comes to managing aPAP."

- Eric

TO HEAR THESE PATIENTS' STORIES, PLEASE VISIT WWW.SAVARAPHARMA.COM

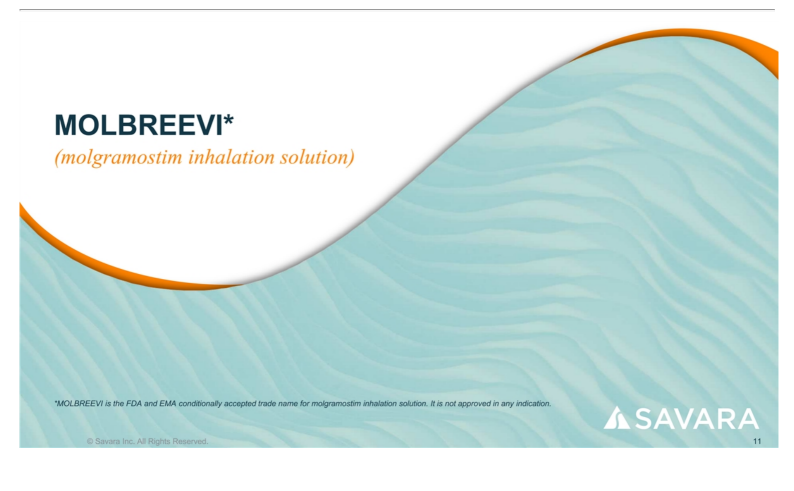
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Disease Burden: Journey of an aPAP Patient

CURRENT JOURNEY POTENTIAL FUTURE JOURNEY Symptoms and Experience Diagnosis with aPAP **Treatment Before Diagnosis** 12 MONTHS 18 MONTHS aPAP ClearPath FROM FIRST EXPERIENCING **AVERAGE TIME FROM FIRST** · Non-invasive, no-cost, simple SYMPTOMS TO SEEING A **SEEING A PHYSICIAN TO** auto-antibody blood test to **DIAGNOSIS PHYSICIAN** help decrease time-todiagnosis aPAP diagnosis Whole lung lavage Insidious development (WLL) MOLBREEVI* Battery of diagnostic tests Potential off-label Typically misdiagnosed as · Clinically meaningful positive therapies pneumonia coupled with Phase 3 results that suggest MOLBREEVI may address the incorrect treatment pathophysiology of aPAP Cycles of misdiagnosis for · Favorable benefit/risk profile months to years · Well tolerated Eventual referral to a pulmonologist for full Rolling BLA initiated pulmonary work-up

*MOLBREEVI is the FDA and EMA conditionally accepted trade name for molgramostim inhalation solution. It is not approved in any indication.

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Savara Investigational Drug-Device Treatment for aPAP

- Once daily 300 μg inhaled MOLBREEVI (inhaled biologic)
- Proprietary eFlow® Nebulizer System (PARI)
 - Optimized for MOLBREEVI administration
 - Well-established manufacturer of devices used for inhalation therapy
 - 5 FDA approved nebulizers based on eFlow® Technology

Nebulization Time: ~5 minutes



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Summary of IMPALA-2 Results

PRIMARY ENDPOINT (MOLBREEVI vs placebo)

0

Change from baseline to Week 24 in DLco% (p=0.0007)1

SECONDARY ENDPOINTS (MOLBREEVI vs placebo)

- Change from baseline to Week 48 in DLco% (p=0.0008)¹
- Change from baseline to Week 24 in SGRQ Total Score (p=0.0072)¹
- Change from baseline to Week 24 in SGRQ Activity Score (p=0.0149)²
- Change from baseline to Week 48 in Exercise Capacity (p=0.0234)²

SAFETY and TOLERABILITY

Well-tolerated; low treatment discontinuation rate (3%), none due to drug-related adverse events

100% of patients who completed the double-blind period enrolled into the open-label period

DLco%, hemoglobin-adjusted percent predicted diffusing capacity of the lungs for carbon monoxide; SGRQ, St. Georges Respiratory Questionnaire iStatistically significant.*Nominally significant.

Phase 3 IMPALA-2 Trial Design

Period 1: Double-blind Period 2: Open-label (top line) (ongoing, not part of top line results) 6-Week Screening n=81 DLCO ≤70% predicted MOLBREEVI 300 μg daily dosing at first screening and baseline MOLBREEVI 300 µg daily dosing Change in % predicted DLCO <15% points to ensure stably impaired W144 placebo patients BL = Primary efficacy analyses (W24) W48 = Durability of efficacy / safety

PRIMARY ENDPOINT

Change from baseline in DLCO at W24

SECONDARY ENDPOINTS

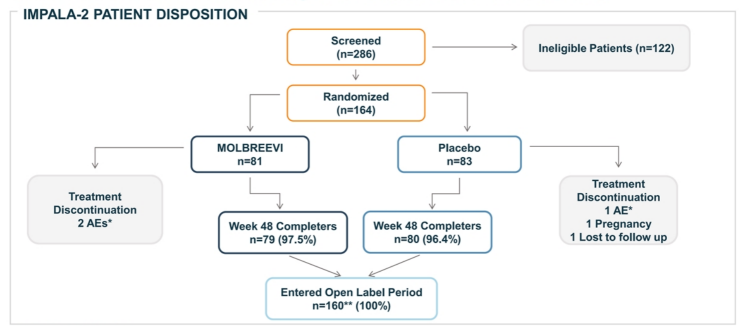
Change from baseline in:

- DLCO at W48
- SGRQ Total Score at W24 and W48
- SGRQ Activity Score at W24 and W48
- Exercise Capacity at W24 and 48

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Discontinuations in Double-Blind Period Were Low: 3%

Participation in Open Label Period Was High: 100% of Double-Blind Period Completers



^{*}Not considered trial drug related
**One placebo patient stopped blinded trial drug but continued trial participation through Week 48 and entered the open label period

Demographics Were Well-Balanced Across Treatment Groups

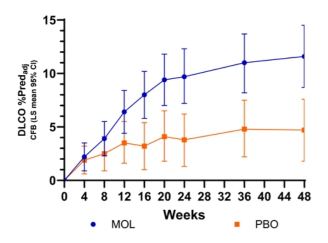
| | | MOLBREEVI N=81 | Placebo N=83 |
|---------------------------|--|--|--|
| Age years | Mean (SD) | 50.8 (13.03) | 48.4 (12.69) |
| Sex n (%) | Male Female | 44 (54.3) 37 (45.7) | 54 (65.1) 29 (34.9) |
| Race n (%) | White Asian Black or African American Other | 38 (46.9) 36 (44.4) 3 (3.7) 4 (4.9) | 40 (48.2) 37 (44.6) 2 (2.4) 4 (4.8) |
| DLCO at baseline | Mean (SD) | 52.6 (11.71) | 52.6 (10.39) |
| DLCO stratification group | ≤ 50% > 50% | 31 (38.3) 50 (61.7) | 32 (38.6) 51 (61.4) |

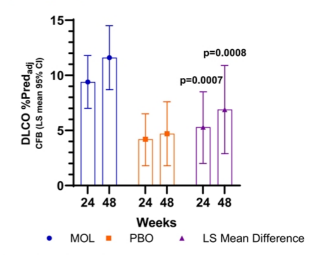
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Primary Endpoint Met (DLCO): Achieved Statistical Significance

MOLBREEVI Superior to Placebo on Change From Baseline in DLCO at W24 (Primary Endpoint) and W48 (Secondary Endpoint)

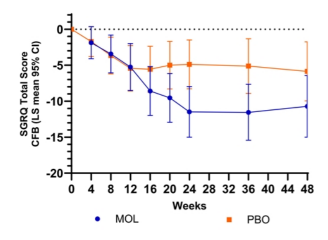


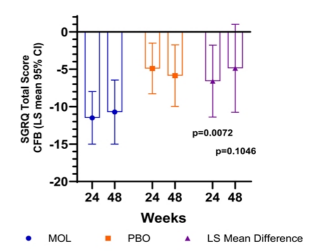


P-values are for difference in LS Mean CFB between MOLBREEVI and placebo

DLCO minimal clinically important difference (MCID) in change from baseline in severe COPD is a 10% increase. MOLBREEVI in aPAP showed a ~10% increase in change from baseline at W24 and ~12% increase in change from baseline at W48.

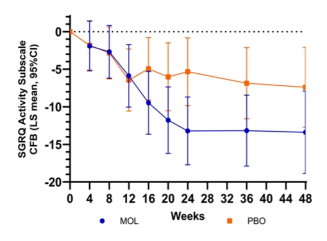
MOLBREEVI Superior to Placebo on Change From Baseline in SGRQ Total Score at W24, Favorability Continues Through W48

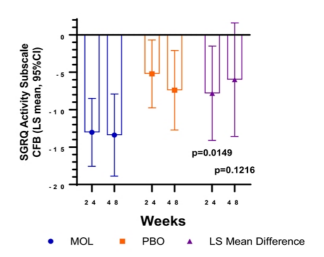




P-values are for difference in LS Mean CFB between MOLBREEVI and placebo

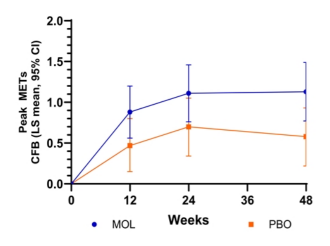
MOLBREEVI Nominally Significant on Change From Baseline in SGRQ Activity Score at W24, Favorability Continues Through W48

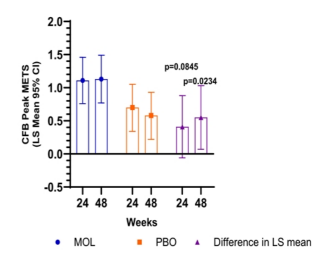




P-values are for difference in LS Mean CFB between MOLBREEVI and placebo

MOLBREEVI Nominally Significant on Change From Baseline in Exercise Capacity (Peak METs) at W48





P-values are for difference in LS Mean CFB between MOLBREEVI and placebo

Lung Lavage Was Permitted as a Rescue Therapy During the Trial

During 48-week double-blind period

- 17 (~10%) patients underwent at least one lung lavage
 - MOLBREEVI: n=6 (7.4%)
 - Placebo: n=11 (13.3%)

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IMPALA-2 Safety Summary: MOLBREEVI Was Well-Tolerated

| Treatment Emergent Adverse Events | MOLBREEVI N=81 n (%) | Placebo N=83 n (%) |
|---|----------------------------|--------------------------|
| Any | 69 (85) | 71 (86) |
| Severe | 13 (16) | 16 (19) |
| Treatment related | 20 (25) | 16 (19) |
| Serious | 14 (17) | 20 (24) |
| Not treatment related | 13 (16) | 20 (24) |
| Treatment related ¹ | 1 (1) | 0 |
| Leading to death | 0 | 0 |
| Leading to trial drug discontinuation | 2 (2) | 1 (1) |
| Special interest (chest pain, hypersensitivity) | 9 (11) | 6 (7) |
| Serious and of special interest | 0 | 1 (1) |

¹SAE of delusions resulting in psychiatric hospitalization in patient with a past medical history of seizure disorder treated with levetiracetam, which is labeled for psychiatric side effects, including delusions; the event was assessed as possibly related to study drug by the investigator.

IMPALA-2 Safety Summary: Most Common Adverse Events

ADVERSE EVENTS IN >10% OF PATIENTS IN ANY TREATMENT ARM DURING DOUBLE-BLIND TREATMENT PERIOD

| Treatment Emergent Adverse Events | MOLBREEVI (N=81) n (%) | Placebo (N=83) n (%) |
|-----------------------------------|------------------------------|----------------------------|
| Any | 69 (85) | 71 (86) |
| Most common | | |
| COVID-19 | 18 (22) | 8 (10) |
| Cough | 17 (21) | 18 (22) |
| Pyrexia | 11 (14) | 9 (11) |
| Nasopharyngitis | 11 (14) | 7 (8) |
| Arthralgia | 9 (11) | 7 (8) |
| Headache | 9 (11) | 7 (8) |
| Diarrhea | 9 (11) | 2 (2) |
| Alveolar proteinosis | 4 (5) | 12 (14) |
| Treatment related | 20 (25) | 16 (19) |

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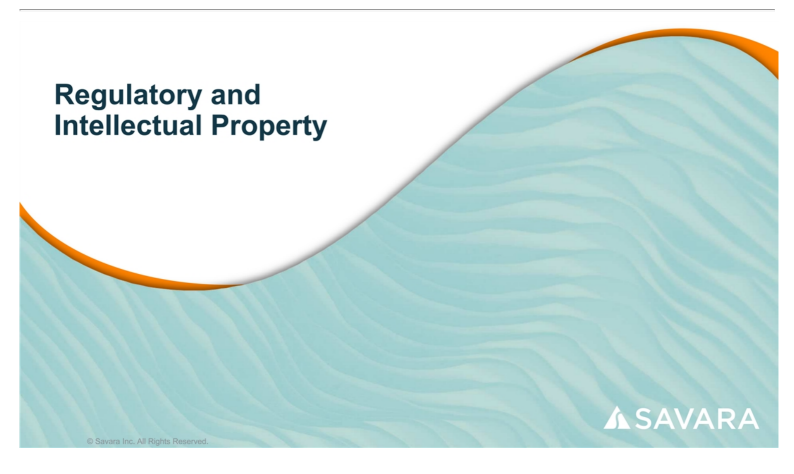
Overview of IMPALA-2 Results: Top Line, DSS, Responder Analyses, and GGO Data

| | Measure | Timeframe | P-Value / Results |
|----------------------------|---------------------------------|--------------------|--|
| | DLco% | Week 24 Week 48 | 0.0007 0.0008 |
| Pulmonary gas exchange | Disease Severity Score (DSS) | Week 24 Week 48 | 0.0239* 0.0006* |
| | Responder Analysis - DLco% | Weeks 24 and 48 | Significantly higher proportions of patients achieved each responder threshold (5%, 7%,10%) with MOLBREEVI compared to placebo |
| | SGRQ Total Score | Week 24 Week 48 | 0.0072 0.1046 |
| Respiratory health-related | SGRQ Activity Score | Week 24 Week 48 | 0.0149 [†] 0.1216 |
| quality of life | Responder Analysis – SGRQ Total | Week 24 Week 48 | Numerically (W24) & significantly (W48) higher proportions of patients achieved each responder threshold (-4, -8, -12-points) with MOLBREEVI compared to placebo |
| Patient functionality | Exercise Capacity (Peak METs) | Week 24 Week 48 | 0.0845 0.0234 † |
| | Chest Computed Tomography – GGO | Week 24 | 0.0004* |
| Surfactant burden | Whole Lung Lavage | Over 48 Weeks | Numerically favorable to MOLBREEVI compared to placebo |

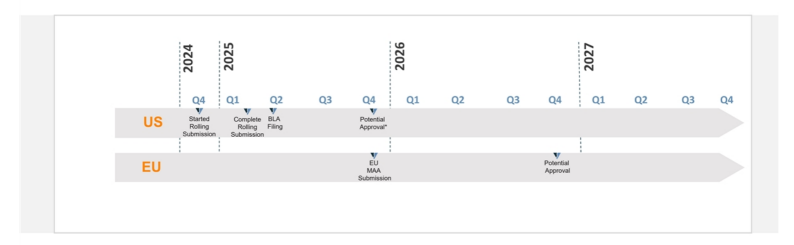
^{*}Post-hoc analysis, 1P-value nominally significant: P-value ≤ 0.0500 but did not meet the p-value threshold required in the pre-specified hierarchical testing procedure.

DLco%, hemoglobin-adjusted percent predicted diffusing capacity of the lungs for carbon monoxide; GGO, ground glass opacification; METs, metabolic equivalents; SGRQ, St. George's Respiratory Questionnaire.

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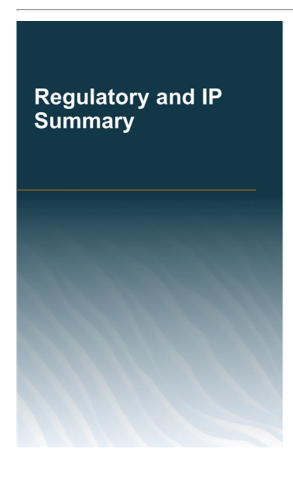


U.S. and European Regulatory Timeline



*Assumes Priority Review is granted by the FDA

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MOLBREEVI IN aPAP REGULATORY DESIGNATIONS

- Orphan Drug Designation, Europe (eligible for 10 years exclusivity)
- Orphan Drug Designation, U.S. (eligible for 7 years exclusivity)
- Fast Track Designation, U.S.
- Breakthrough Therapy Designation, U.S.
- Innovation Passport Designation, U.K.
- Promising Innovative Medicine Designation, U.K.

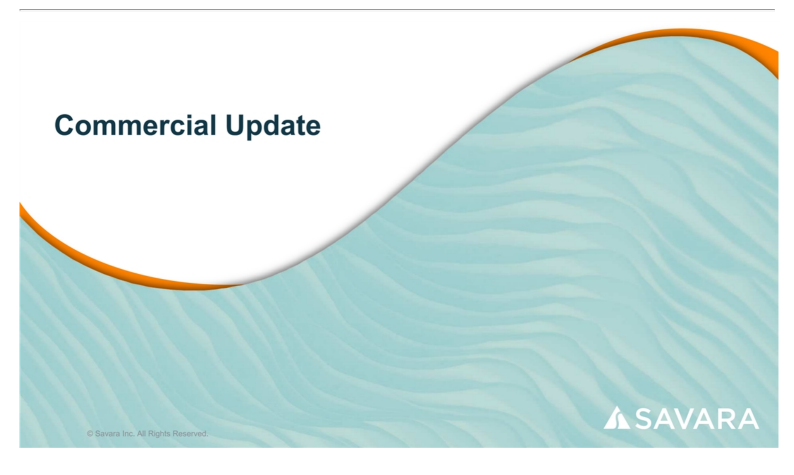
BIOLOGIC EXCLUSIVITY

 Upon Biologics License Application (BLA) approval FDA would grant 12 years marketing exclusivity

INTELLECTUAL PROPERTY

- Pending patent applications for MOLBREEVI drug formulation and methods of use including treating aPAP with MOLBREEVI
- Worldwide exclusive license to proprietary eFlow[®] Nebulizer System (PARI) for MOLBREEVI in aPAP and pending joint patent application with PARI for the drug/device combination
- Proprietary cell bank for MOLBREEVI

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Commercial Launch Planning Advancing Against Near Term Objectives

AWARENESS

Expand awareness of autoimmune PAP among targeted health care professional and patients



INFRASTRUCTURE

Build critical capabilities to facilitate access to MOLBREEVI post approval



PERSONNEL

Hire and onboard key commercial roles to expand core activities



TESTING

Evolve antibody testing platform with an eye toward long term market expansion

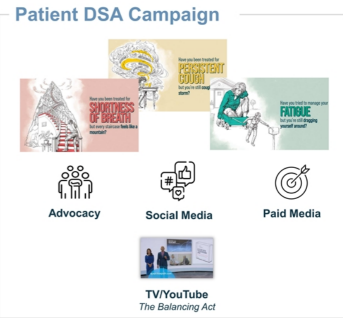


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Autoimmune PAP Disease State Awareness Campaign

Multi-channel effort across healthcare professionals and patients





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Exclusive Specialty Pharmacy with Integrated Patient Services Right-sized model for first-to-market solution for orphan condition



- Smaller patient population is best served by a single specialty pharmacy
 - Consistency
 - · Seamless provision of services
 - Clear visibility to all patient data to inform key performance indicators
- Currently evaluating partners
 - Relevant pulmonary experience
 - Demonstrated track record of exceptional patient and provider services

Single source pharmacy will service all patients with direct shipments and ongoing support

MyMolbreevi: Best in Class Support Program in Development Program aims to reduce access barriers for appropriate MOLBREEVI patients post approval



PATIENT SERVICES

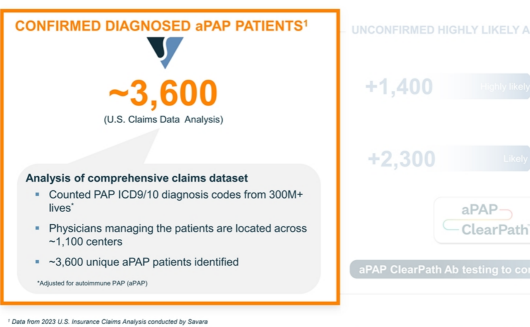
- Case management approach
 - Dedicated care navigator
 - Dedicated care name
 Single point of contact
- Financial assistance
 - Commercial co-pay program
 - Free drug for eligible patients
- Clinical education
 - Pharmacist calls
 - Device training
 - Nurse educators
 - Adherence support
- Insurance services
 - Prior authorization
 - Appeals



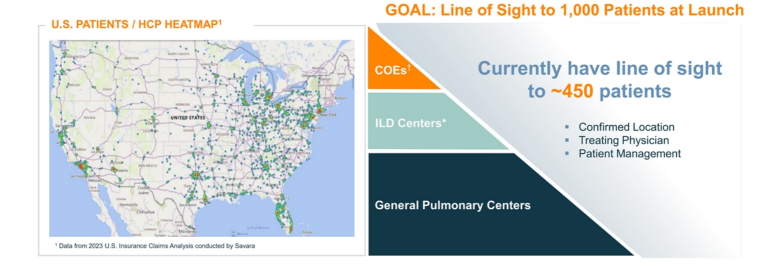
PRESCRIBER SERVICES

- Streamlined prescribing
- Prior authorization checklist
- Sample letter of medical necessity
- Sample letter of appeal

Significant U.S. Opportunity with ~3,600 Currently Diagnosed aPAP Patients



U.S. Centers Prioritized Based on Experience Treating aPAP Patients



PAP Center of Excellence (COE) includes healthcare organizations listed on PAP Foundation website, IMPALA-2 clinical trial sites, and other sites of expertise ILD clinics are dedicated to the management of patients with a wide variety of interstitial lung diseases that can range from pulmonary fibrosis to rare lung diseases

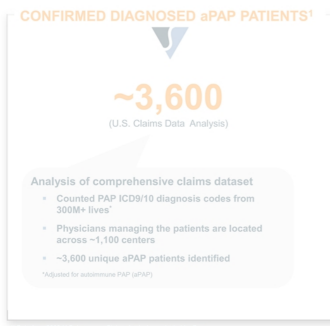
Market Development Team

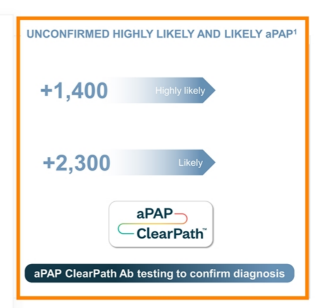
Broadens market reach and accelerates pre-launch activities



- Projected market development team of ~25 people (including leadership)
- Target list of accounts expanded beyond current field medical list to broaden reach
- Territory managers will be added in waves gated to key milestones
- Key activities include:
 - Profiling accounts to gain line of sight into currently diagnosed patients
 - aPAP disease awareness and education
 - Dry blood spot (DBS) antibody testing education

Claims Data Analysis Suggests U.S. Market May Be 2x Larger





1) Data from 2023 U.S. Insurance Claims Analysis conducted by Savara

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aPAP ClearPath Testing Platform

No cost antibody testing to identify aPAP among undiagnosed patients

CURRENTLY AVAILABLE: SERUM





- Launched in the U.S. and Europe
- Platform used in Interstitial Lung Disease (ILD) Clinic Pilot Program





- Simple finger prick performed in a physician's office
- Removes logistical challenges to serum testing
- Target launch end of 1Q 2025

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Europe (EU4+UK) Market Development is Underway

TREATMENT CENTER MAPPING¹



| Country | Key Centers ¹ | Est. TAM ² |
|---------|--------------------------|-----------------------|
| Germany | 11 | ~1,100 |
| UK | 25 | ~900 |
| France | 24 | ~900 |
| Italy | 16 | ~700 |
| Spain | 12 | ~600 |
| Total | 88 | ~5,000 |

- aPAP Centers of Excellence identified (8)
- aPAP ClearPath antibody test launched in Europe
- 62 patients in Europe enrolled in IMPALA-2 trial open-label extension³

¹ Savara 2024 EU4+ UK Primary (N= 6 EU4+ UK Principal Investigators, 5 EU4+UK Lab Directors) and Secondary Market Research ² Data from 2023 U.S. Insurance Claims Analysis conducted by Savara and extrapolated based on geographic population ³ Enrolled across 15 IMPALA-2 sites in the EU, UK, and Turkey

MOLBREEVI: U.S. Commercial Opportunity

MOLBREEVI

- Clinically meaningful benefit
- Strong stakeholder interest
- Orphan drug pricing potential
- Chronic dosing

Long Term Exclusivity

- 12-year biologic exclusivity (U.S.)
- Biosimilar competition unlikely



Efficient Rare Disease Model

- Small customer facing footprint
- Exclusive pharmacy network

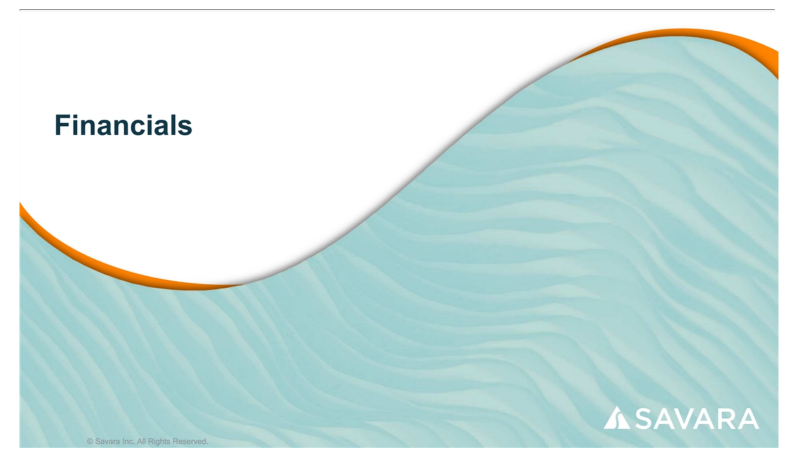
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Significant Unmet Need

No FDA approved therapies

High disease burden

 Whole lung lavage is invasive and not standardized



- Cash runway through 2Q 2027
 - ~\$219M in cash, cash equivalents and short-term investments*
- Strong investor support with coverage from 8 equity research analysts

ANALYST COVERAGE

| Jefferies | Andrew Tsai |
|-----------------------|-----------------------|
| Piper Sandler | Yasmeen Rahimi, PhD |
| Guggenheim Securities | Vamil Divan, MD, MBA |
| Oppenheimer | Francois Brisebois |
| JMP | Jonathan Wolleben |
| H.C. Wainwright | Andrew Fein |
| Evercore ISI | Liisa Bayko, MSC, MBA |
| Wells Fargo | Tiago Fauth |
| | |

*As of 9/30/24



Near- and Long-Term U.S. Market Opportunity in aPAP is Sizeable

~3,600 Current U.S. TAM of confirmed diagnosed patients
 \$300K-\$500K Orphan rare disease potential pricing power
 ~3,700 Large pool of likely patients that are currently undiagnosed
 Multiple Patents currently being prosecuted
 12-years Biologic exclusivity in U.S. upon approval

Long-term Durable revenue stream with biosimilar competition unlikely



TAM = Total addressable market

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THANK YOU