

Developing New Therapies for Rare Respiratory Diseases

December 2024



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Executive Leadership Team

Matthew Pauls, J.D., M.B.A.

Chair & Chief Executive Officer

Anne Erickson
Chief Business Officer

Dave Lowrance
Chief Financial &
Administrative Officer

Rob Lutz, M.B.A.

Chief Operating Officer

Braden Parker, M.B.A.

Chief Commercial Officer

Ray Pratt, M.D. FACP Chief Medical Officer

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EVP, Global Technical

Operations

Yasmine Wasfi, M.D., Ph.D. EVP, Head of Clinical Operations/Development



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





aPAP and MOLBREEVI* (molgramostim inhalation solution)

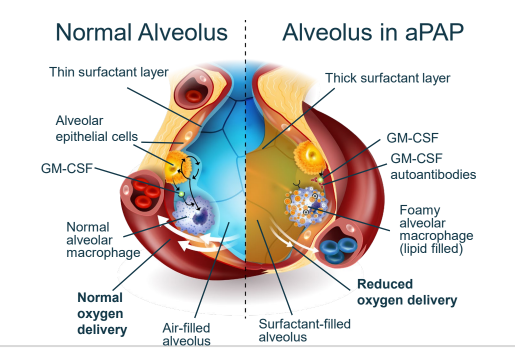


aPAP: An Autoimmune Disease of Alveolar Macrophage Dysfunction

NORMAL LUNG FUNCTION

Alveoli need surfactant to keep from collapsing

GM-CSF is critical for alveolar macrophage function and allows for alveolar surfactant homeostasis, structure, function, and host defense

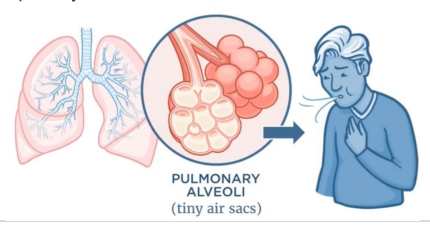




Rare lung disease caused by GM-CSF autoantibodies which block GM-CSF signaling and reduce surfactant clearance. This results in:

Surfactant accumulation that blocks movement of oxygen from the alveoli into the blood

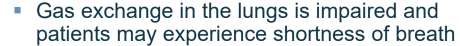
Reduced blood oxygenation results in difficulty breathing and, ultimately, hypoxemic respiratory failure





aPAP is a Rare, Long-Term, Chronic Disease

Progressive Shortness of Breath



 At first it occurs upon exertion, but as disease progresses, it can occur even when a person is at rest

Cough and Episodes of Fever

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

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

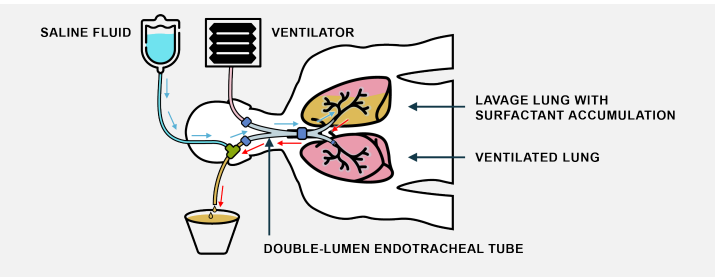
 In the long-term, the disease can lead to serious complications, including fibrosis, and may lead to the need for lung transplantation



There are no approved drugs for the treatment of aPAP. Only option is a lung lavage, an invasive procedure.



- A lung lavage physically removes excess surfactant from the lungs and requires hospitalization
- Performed under general anesthesia
- Unavailable at many medical institutions



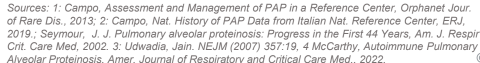
A Lung Lavage is an Invasive Procedure Performed in a Tertiary Center and is Not Standardized



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

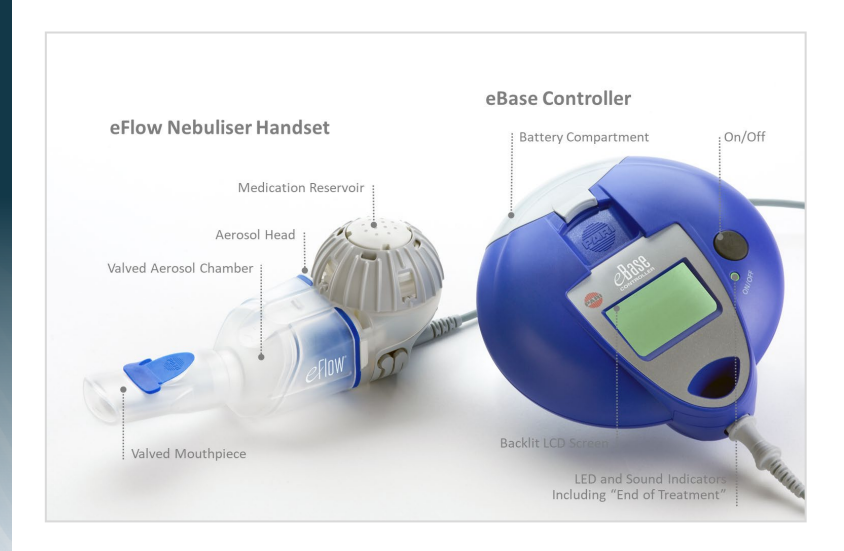
Saline is drained by gravity and continued until lavage fluid becomes clear





Savara Investigational Drug-Device Treatment for aPAP

- Once daily 300 μg inhaled MOLBREEVI
- 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



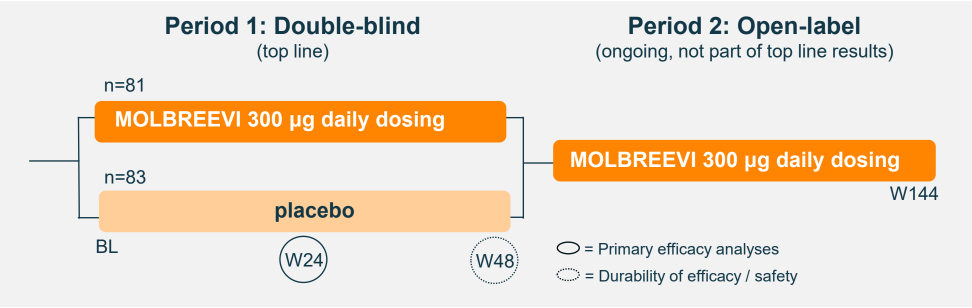


Phase 3 IMPALA-2 Top Line Results

Phase 3 IMPALA-2 Trial Design

6-Week Screening

- DLCO ≤70% predicted at first screening and baseline
- Change in % predicted DLCO <15% points to ensure stably impaired patients



PRIMARY ENDPOINT

Change from baseline in DLCO at W24

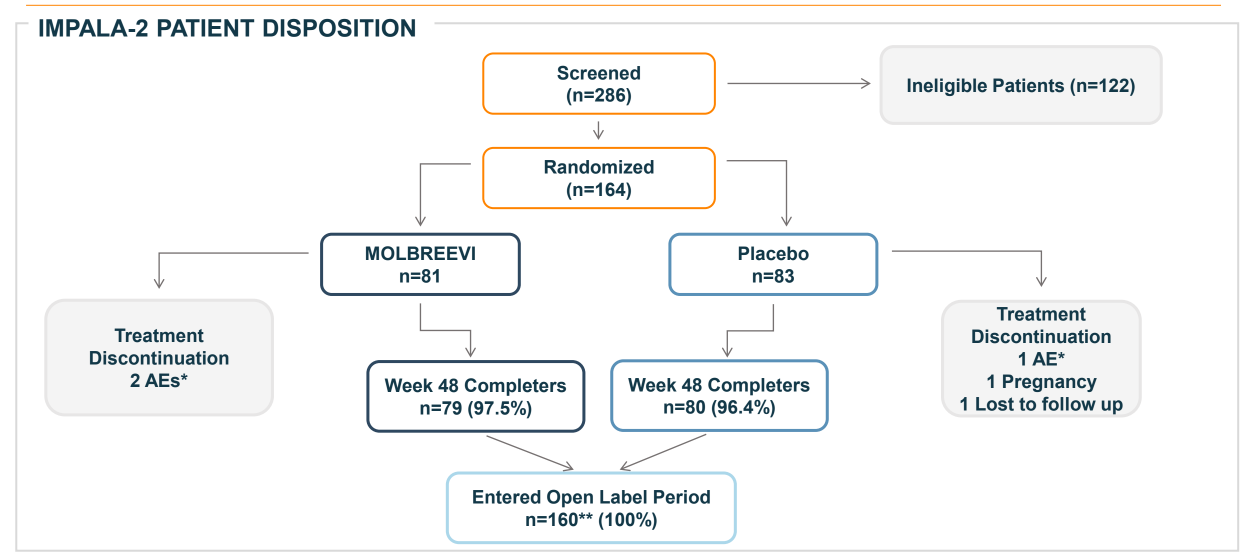
SECONDARY ENDPOINTS

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



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 © Savara Inc. All Rights Reserved.

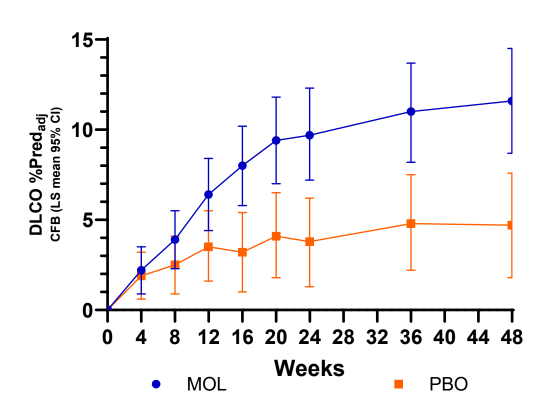
Demographics Were Well-Balanced Across Treatment Groups

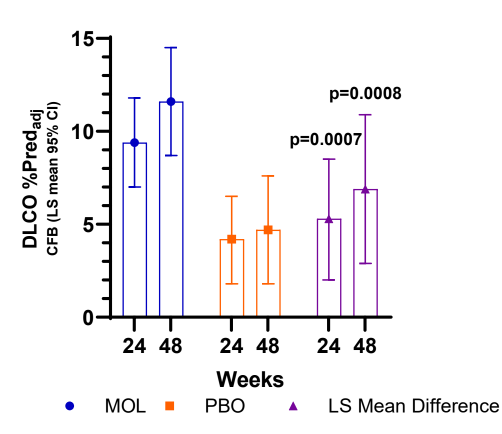
		MOLBREEVI N=81	Placebo N=83
Age years	Mean (SD)	50.8 (13.03)	48.4 (12.69)
Sex n (%)	Male Female	44 (54.3) 37 (45.7)	54 (65.1) 29 (34.9)
Race n (%)	White Asian Black or African American Other	38 (46.9) 36 (44.4) 3 (3.7) 4 (4.9)	40 (48.2) 37 (44.6) 2 (2.4) 4 (4.8)
DLCO at baseline	Mean (SD)	52.6 (11.71)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



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)

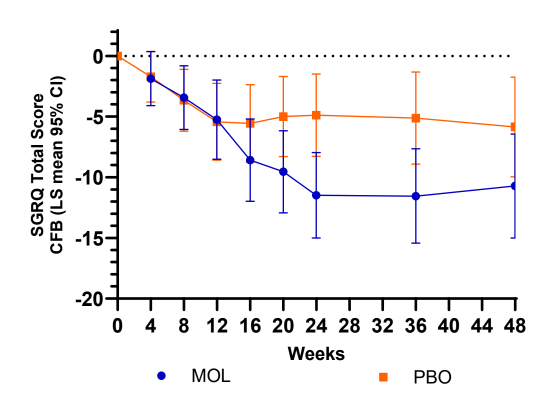


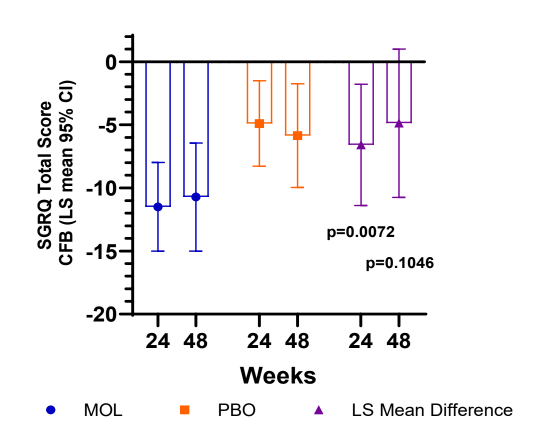


P-values are for difference in LS Mean CFB between MOLBREEVI and placebo

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. SAVARA

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

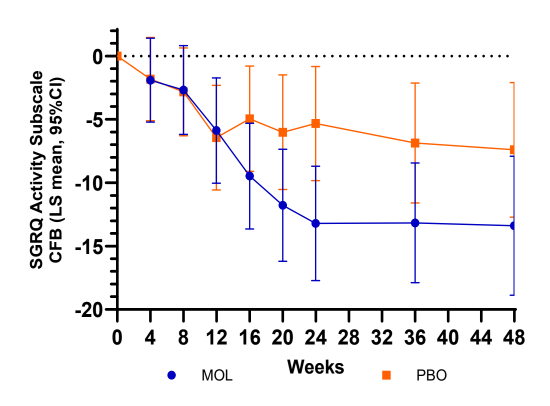


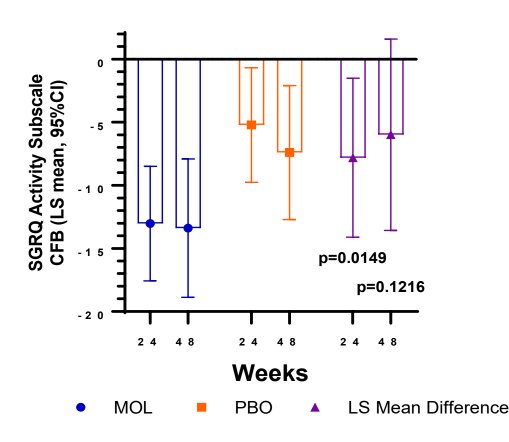


P-values are for difference in LS Mean CFB between MOLBREEVI and placebo



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

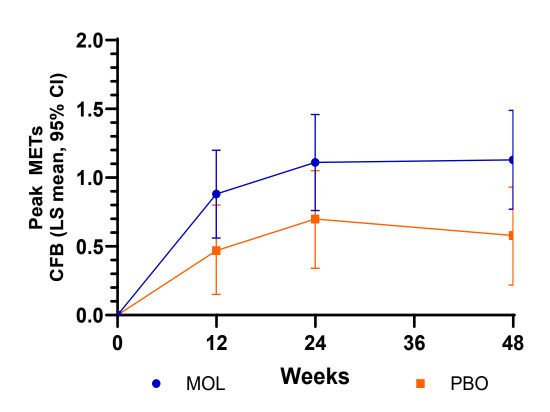


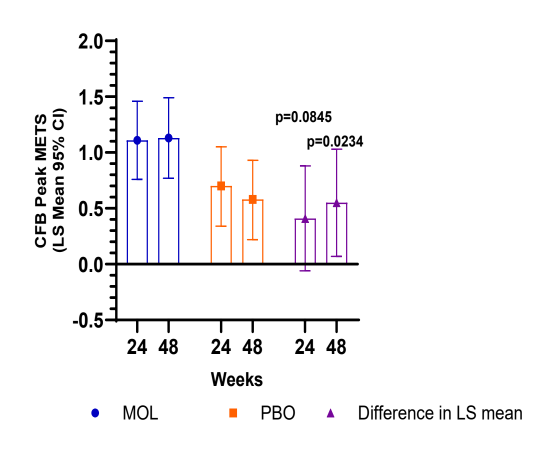


P-values are for difference in LS Mean CFB between MOLBREEVI and placebo



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%)



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)



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. †P-value nominally significant: P-value ≤ 0.0500 but did not meet the p-value threshold required in the pre-specified hierarchical testing procedure.



Regulatory and Intellectual Property

United States: BLA Rolling Submission Initiated in Dec. 2024, on Track to be Completed by End of 1Q 2025

Europe: MAA Submission on Track for End of 2025

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.

IMPALA-2

 Trial design endorsed by regulatory authorities in the U.S., Canada, Japan, South Korea, Australia, U.K., and countries in Europe where the trial is being conducted

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



Commercial Outlook



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

CONFIRMED DIAGNOSED aPAP PATIENTS¹



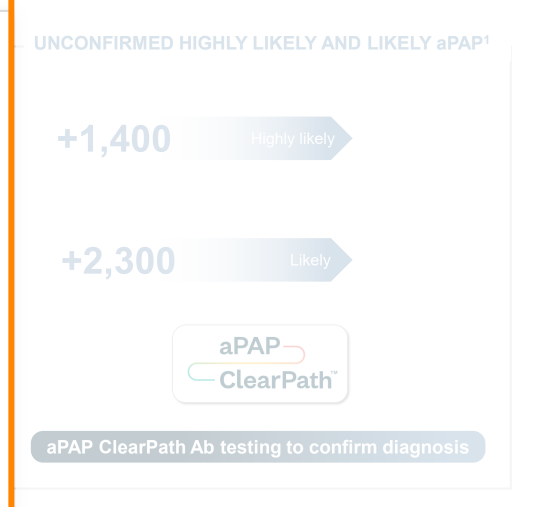
~3,600

(U.S. Claims Data Analysis)

Analysis of comprehensive claims dataset

- Counted PAP ICD9/10 diagnosis codes from 300M+ lives*
- Physicians managing the patients are located across ~1,100 centers
- ~3,600 unique aPAP patients identified

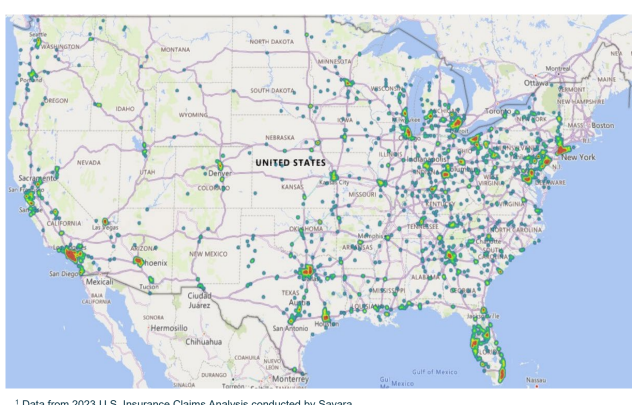
*Adjusted for autoimmune PAP (aPAP)



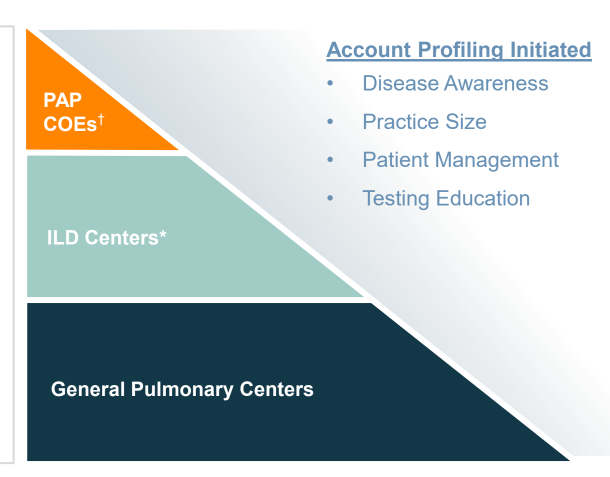


U.S. Centers Prioritized Based on Experience Treating aPAP Patients

U.S. PATIENTS / HCP HEATMAP¹



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





Overwhelming Support for MOLBREEVI Across Stakeholders

U.S. PULMONOLOGISTS

83%

Likely to prescribe MOLBREEVI regardless of severity of disease

Perceived MOLBREEVI as superior to current options based on:

- Overall efficacy
- Simple, standard nebulizer
- Favorable safety profile
- FDA approval

U.S. PAYERS

87%

Intend to cover MOLBREEVI with typical Prior Authorization criteria when priced between \$300-500K annually

Not concerned about impact on pharmacy budget

Recognize the significant disease burden associated with aPAP

U.S. PATIENTS

100%

Think new, non-invasive PAP treatments are needed

WOULD ask their doctor to prescribe MOLBREEVI

WOULD take MOLBREEVI if their doctor recommended it

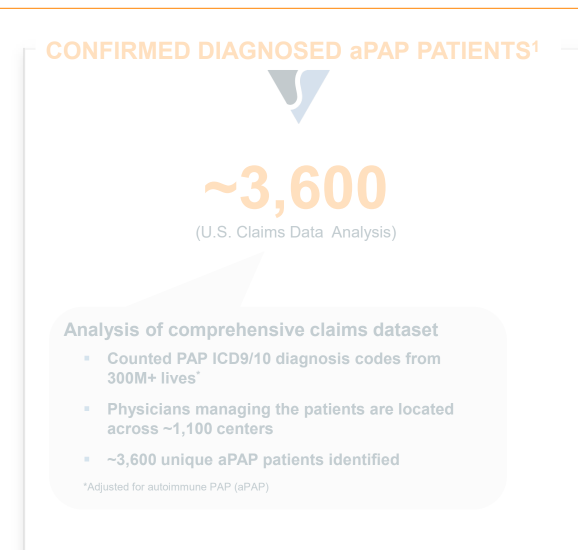
Savara 2024 Qualitative Research: N=10 US KOLs and high-volume pulmonologists; Quantitative research: N=78 HCPs

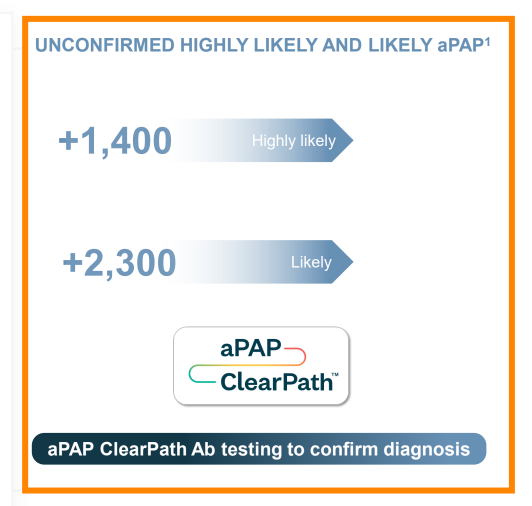
Savara 2024 U.S. payer pricing and reimbursement research: N=10 representing ~88M covered lives

Savara 2024 Patient Advisory Board meetings: N = 7 aPAP patients



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







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 expected to launch in Europe by end of 2024 62 EU patients enrolled in IMPALA-2 open label extension study*



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: Commercial Opportunity

Significant Unmet Need

- High disease burden
- No FDA approved therapies
- Whole lung lavage is invasive, not standardized, and minimally used

Efficient Rare Disease Model

- Small customer facing footprint
- Exclusive pharmacy network
- Regional expansion optionality (go-it-alone, partnership, etc.)



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



Financials



- 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
JMP H.C. Wainwright	Jonathan Wolleben Andrew Fein

*As of 9/30/24

Financial Highlights



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Thank You

