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

FORM 8-K

CURRENT REPORT

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

Date of Report (Date of earliest event reported): December 11, 2024

RESHAPE LIFESCIENCES INC.

(Exact name of registrant as specified in its charter)

1-37897 (Commission File Number) 26-1828101 (I.R.S. Employer Identification Number)

18 Technology Drive, Suite 110,

Irvine, CA (Address of principal executive offices)

(949) 429-6680

(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)

Delaware (State or other jurisdiction of

incorporation)

□ Soliciting material pursuant to Rule 14a-12 under the Exchange Act (17 CFR 240.14a-12)

D Pre-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))

D 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 Exchange on which Registered The Nasdaq Capital Market Title of Class Symbol Common stock, \$0.001 par value per share RSLS

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 \Box

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

92618 (Zip Code)

Item 7.01 Regulation FD Disclosure.

Representatives of ReShape Lifesciences Inc. (the "Company") and Vyome Therapeutics, Inc. intend to make presentations at investor conferences and in other forums and these presentations may include the information contained in Exhibit 99.1 attached to this Current Report on Form 8-K. A copy of the presentation slides containing such information that may be disclosed by the Company is attached as Exhibit 99.1 to this report and the information set forth therein is incorporated herein by reference and constitutes a part of this report.

The Company is furnishing the information contained in Exhibit 99.1 pursuant to Regulation FD and Item 7.01 of Form 8-K promulgated by the Securities and Exchange Commission ("SEC"). This information shall not be deemed to be "filed" with the SEC 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, nor shall it be deemed to be incorporated by reference in any filing under the Securities Act of 1933, as amended, or the Exchange Act, except as expressly set forth by specific reference in such filing.

The information contained in Exhibit 99.1 is summary information that is intended to be considered in the context of the Company's SEC filings and other public announcements that the Company may make, by press release or otherwise, from time to time. The Company undertakes no duty or obligation to publicly update or revise the information contained in Exhibit 99.1, although it may do so from time to time as its management believes is warranted. Any such updating may be made through the filing of other reports or documents with the SEC, through press releases or through other public disclosure. By filing this report and furnishing this information, the Company makes no admission as to the materiality of any information contained in this report, including Exhibit 99.1.

Additional Information

In connection with the proposed merger with Vyome Therapeutics, Inc. (the "Merge") and sale of assets to Biorad Medisys of an affiliate thereof (the "Asset Sale"), ReShape plans to file with the Securities and Exchange Commission (the "SEC") and mail or otherwise provide to its stockholders a joint proxy statement/prospectus and other relevant documents in connection with the proposed Merger and Asset Sale. Before making a voting decision, ReShape's stockholders are urged to read the joint proxy statement/prospectus and other relevant documents in connection with the proposed Merger and Asset Sale or incorporated by reference therein carefully and in their entirety when they become available because they will contain important information about ReShape, Vyome and the proposed transactions. Investors and stockholders may obtain a free copy of these materials (when they are available) and other documents filed by ReShape at the SEC is website at www.sec.gov, at ReShape's website at www.reshapelifesciences.com, or by sending a written request to ReShape at 18 Technology Drive, Suite 110, Irvine, California 92618, Attention: Corporate Secretary.

Participants in the Solicitation

This document does not constitute a solicitation of proxy, an offer to purchase or a solicitation of an offer to sell any securities of ReShape and its directors, executive officers and certain other members of management and employees may be deemed to be participants in soliciting proxies from its stockholders in connection with the proposed Merger and Asset Sale. Information regarding the persons who may, under the rules of the SEC, be considered to be participants in the solicitation of ReShape's stockholders in connection with the proposed Merger and Asset Sale will be set forth in joint proxy statement/prospectus if and when it is filed with the SEC by ReShape and Vyome. Security holders may obtain information regarding the names, affiliations and interests of ReShape's directors and officers in ReShape's Annual Report on Form 10-K for the fiscal year ended December 31, 2023, which was filed with the SEC on April 1, 2024. To the extent the holdings of ReShape securities by ReShape's directors and executive officers have changed since the amounts set forth in ReShape's proxy statement for its most recent annual meeting of stockholders, such changes have been or will be reflected on Statements of Change in Ownership on Form 4 filed with the SEC. Additional information regarding these individuals and any direct or indirect interests they may have in the proposed Merger and Asset Sale will be set forth in the joint proxy statement/prospectus when and if it is filed with the SEC in connection with the proposed Merger and Asset Sale, at ReShape's website at www.reshapelifesciences.com.

Forward-Looking Statements

Certain statements contained in this filing may be considered forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995, including statements regarding the Merger and Asset Sale and the ability to consummate the Merger and Asset Sale. These forward-looking statements generally include statements that are predictive in nature and depend upon or refer to future events or conditions, and include words such as "believes," "plans," "anticipates," "estimates," "expects," "intends," "strategy," "future," "opportunity," "may," "will," "should," "potential," or similar expressions. Statements that are not historical facts are forward-looking statements. Forward-looking statements beade on current beliefs and assumptions that are subject to risks and uncertainties. Forward-looking statements are based on current beliefs and assumptions that are subject to risks and uncertainties. Forward-looking statements are based on current beliefs and assumptions that are subject to risks and uncertainties. Forward-looking statements are based on current beliefs and assumptions that are subject to risks and uncertainties. Forward-looking statements are a result of various factors, including, without limitation: (1) ReShape may be unable to obtain stockholder approval as required for the proposed Merger and Asset Sale; (2) conditions to the closing of the Merger or Asset Sale may involve unexpected costs, liabilities or delays; (4) ReShape's business and/or competitive factors; (7) the occurrence of any event, change or other circumstances that could give rise to the termination of the Merger and Asset Purchase Agreement (8) the effect of the announcement of the Merger and Asset Sale; (3) the offect of the announcement of the future results of ReShape are set forth in its filings with the SEC, which are available on the SEC's website at www.sec.gov, specifically under the heading "Risk factors." The risks and uncertainties described above and in ReShape's most recent filed Annual Report on F

Item 9.01 Financial Statements and Exhibits.

(d)	Exhibits.

_	Exhibit No.	Description	Method of Filing
_	<u>99.1</u>	Investor Presentation Slides	Furnished herewith
	104	Cover Page Interactive Data File (the cover page XBRL tags are embedded in the Inline XBRL	
		document).	

SIGNATURES

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.

ENTEROMEDICS INC.

By: /s/ Paul F. Hickey Paul F. Hickey Chief Executive Officer

Dated: December 11, 2024





A US-India healthcare platform with clinical-stage immuno-inflammation assets

PENDING MERGER \$RSLS TO \$HIND

CORPORATE PRESENTATION

DECEMBER 2024

VYOME THERAPEUTICS, INC. ("Vyome")

Any statements contained in this presentation that do not describe historical facts may constitute forward-looking statements as that term is defined in the Private Securities Litigation Reform Act of 1995. These statements may be identified by words such as "believe," "expect," "may," "plan," "potential," "will," and similar expressions, and are based on Vyome's current beliefs and expectations. These forward-looking statements include expectations regarding Vyome's development of its drug candidates, including the timing of its clinical trials and regulatory submissions. These statements involve risks and uncertainties that could cause actual results to differ materially from those reflected in such statements. Risks and uncertainties that may cause actual results to differ materially include uncertainties inherent in the conduct of clinical trials, Vyome's reliance on third parties over which it may not always have full control, public health crises, epidemics and pandemics such as the COVID-19 pandemic, including its impact on the timing of Vyome's regulatory and research and development activities, Any forward-looking statements speak only as of the date of this presentation and are based on information available to Vyome as of the date of this presentation, future events or otherwise.

This presentation also contains estimates and other statistical data made by independent parties and by us relating to market size and other data about our industry. These data involve a number of assumptions and limitations, and you are cautioned not to give undue weight to such estimates. In addition, projections, assumptions and estimates of our future performance and the future performance of the markets in which we operate are necessarily subject to a high degree of uncertainty and risk. References to any publications, reports, surveys or articles prepared by third parties should not be construed as depicting the complete findings of the entire publication, report, survey or article. The information in any such publication, report, surveys or article is not incorporated by reference in this presentation.

Vyome is building a 3-pillared healthcare platform in the US-India innovation corridor

.We intend to list on Nasdaq via reverse merger with \$RSLS under the ticker \$HIND

We have invested nearly a decade and millions of dollars to build a set of immuno-inflammatory assets with several 12-24 month catalysts

DID YOU KNOW ...

HIND

an ancient name for India

*Allied Market Research - Anti-Inflammatory Therapeutics Market Review

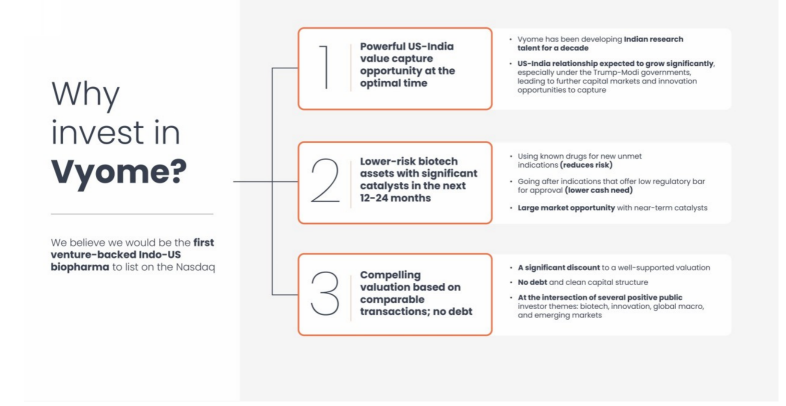
\$125B+ by 2028*

The immuno-inflammatory market is expected to be Cambridge, MA HQ

Home to MIT and Harvard

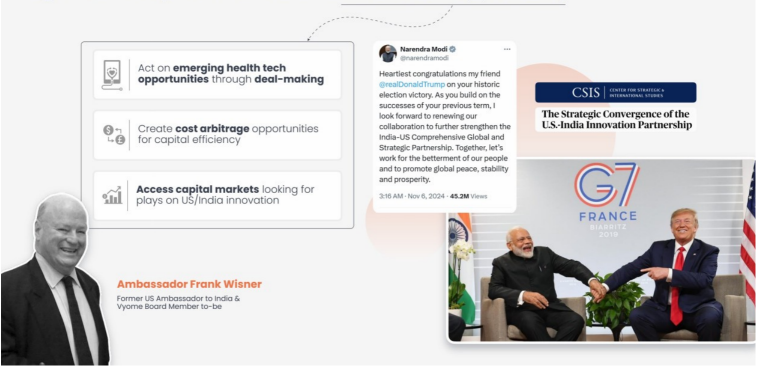
At Vyome, we are passionate about **transforming healthcare** based on world-class science & talent leveraging the **US-India innovation corridor**





US-India special relationship and our positioning unlocks

significant capital markets and innovation opportunity



The current biotech assets offer a low-risk way to unlock significant value over the next 12-24 months

Vyome assets	Market size	Capital required	Potential value inflection timeline
Malignant Fungating Wound	\$2.5B ¹	\$6M	2027 (pivotal data)
Uveitis (steroid replacement)	\$2.6B ² by 2032	\$3M	2026 Phase 1
Inflammatory acne*	\$6.0B ³ by 2028*	-	2025 Potential partnership deal

1 Based on 60-100K patients in U.S. (10% of -600K cancer deaths per year in U.S.), and 1M patients globally (10% of ~10M advanced cancer patients). Based on average of 5-15% incidence of cancer patients developing MFW (from The Microbiome, Malignant Fungating Wounds, and Palliative Care. Front. Cell. Infect. Microbiol., November 2019). 2. https://www.imarcgroup.com/uveitis-treatment-market 3. Total global acce market is forecasted to grow to \$11.6B by 2028. https://www.marketdataforecast.com/market-reports/acne-medication-market Inflammatory acne comprises over 50% of the acne market. https://www.verifiedmarketresearch.com/product/acne-medication-market/

Malignant Fungating Wound (MFW) is \$1B/year unmet need with no approved drug; the latest data on Vyome's drug shows strong efficacy

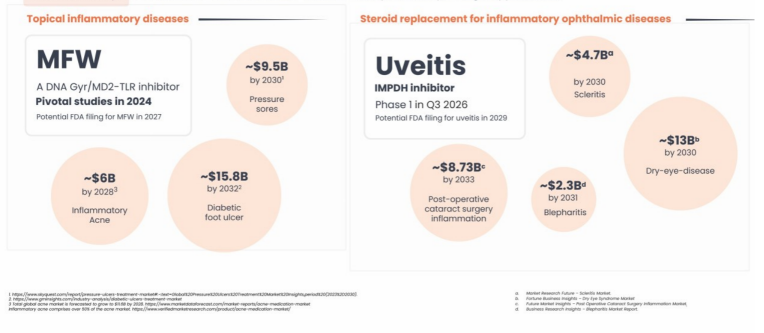


2 The Microbiome, Malignant Fungating Wounds, and Palilative Care. Front. Cell. Infect. Microbial, November 2019, EASED study (2023) https://acsiaumais.on/neilstrarv.wijey.com/doi/10.3322/cage.21820

Vyome's 2 biotech asset areas form a pipeline **addressing large unmet markets**

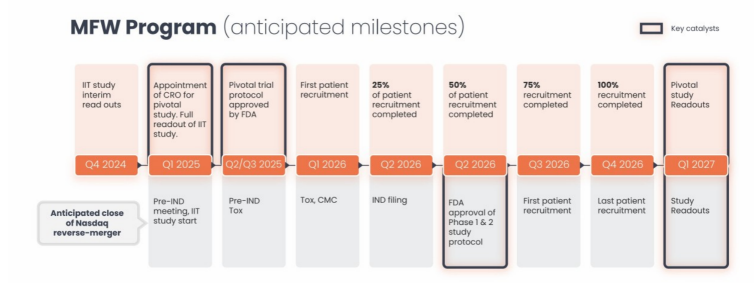
Our strategy

Use an unmet orphan indication for low-risk and cost-efficient development to open larger opportunities



Vyome is focused on unlocking multiple value inflection points

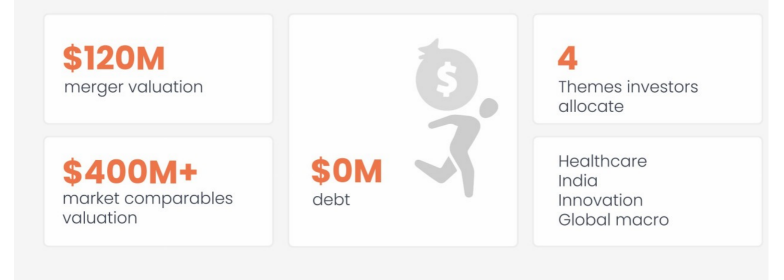
from its core pharma assets over the next 12-24 months



Uveitis Program (anticipated milestones)

Vyome has a compelling entry valuation, no debt,

and a narrative appealing to several types of investors



Public Comparables Inflammation & Immunology Market Players

Average Market Cap. of **\$419M** (not including Immunovant)

	VYOME	INmuneBio	gosametrio	prime TTD	🝐 alumis	UpstreamBIO	WIMMUNOVANT
Lead Indication	Malignant fun gating wound	Metastatic Castrate Resistant Prostate Cancer	Pulmonary Arterial Hypertension	Chronic Granulomatous Disease (CGD)	Psoriasis	Severe Asthma, CRSwNP, & COPD	Myasthenia Gravis & Thyroid Eye Disease
Stage / Phase		Phase 1/2	Phase 2	Phase 1/2	Phase 2	Phase 2	Phase 3
Market Size	~\$3B (Current)	~\$13B (2023)	~\$7B (2023)	~1.3B (2023)	~\$25B (2023)	~7.5B (2023)	~\$4B (2023)
Program Stages		3 clinical 0 pre-clinical	1 clinical 0 pre-clinical	1 clinical 5 pre-clinical	3 clinical 2 pre-clinical	3 clinical 0 pre-clinical	4 clinical 1 pre-clinical
Market Cap (\$MM)		\$134	\$198	\$529	\$534	\$1,271	\$4,314

Denotes aspirational comparable company

Notes:

Precedent M&A Comparables Acquired Inflammation & Immunology Market Players

Average Market Cap. of **\$2.8B** (includes Morphic, DICE, Alpine, Arena, Prometheus) and Transaction Price of **\$482M** (includes Landos and Escient)

	VYOME		escient	MORPHIC	DICC		ARENA	Prometheus
Lead Indication	Malignant fun gating wound	Ulcerative colitis	Atopic dermatitis	Inflammatory bowel disease	Psoriasis	Systemic lupus erythematosus	Ulcerative colitis	Inflammatory bowel disease
Stage / Phase	Phase 2	Phase 2	Phase 2	Phase 2	Phase 2	Phase 2	Phase 3	Phase 2
Market Size	~\$3B (Current)	~\$9B (2031)	~\$9.3B (2023)	~\$30B (Current)	~\$25B (2023)	~\$3B (2025)	~\$9B (2031)	~\$30B (Current)
Program Stages	2 clinical 1 pre-clinical	1 clinical 3 pre-clinical	2 clinical 0 pre-clinical	1 clinical 0 pre-clinical	1 clinical 3 pre-clinical	2 clinical 2 pre-clinical	4 clinical 1 pre-clinical	1 clinical 0 pre-clinical
Market Cap (\$MM)	\$120	\$213 ⁽³⁾	\$750 ^(d)	\$1,595	\$1,616	\$2,553	\$3,063	\$5,419

Denotes private company at time of acquisition

Vyome has an ideal team to execute its plan Deep expertise in building and scaling companies; scientific thought leaders in drug development; extensive US-India crossborder experience



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