

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): **August 13, 2026**

PELTHOS THERAPEUTICS INC.

(Exact name of registrant as specified in its charter)

Nevada

(State or other jurisdiction
of incorporation)

001-41964

(Commission File Number)

86-3335449

(IRS Employer
Identification No.)

4020 Stirrup Creek Drive, Suite 110

Durham, NC

(Address of registrant's principal executive office)

27703

(Zip code)

Registrant's telephone number, including area code: **(919) 908-2400**

(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):

- ☐ 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 \$0.0001 per share	PTHS	The NYSE American LLC

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

Emerging growth company ☒ x

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. ☐ o

Item 2.02. Results of Operations and Financial Condition.

On August 13, 2026, Pelthos Therapeutics Inc. (the “Company”) issued a press release summarizing its financial results for the second quarter 2026, as well as providing an update on the Company’s operations. A copy of the press release is furnished as Exhibit 99.1 to this Current Report on Form 8-K and is incorporated by reference.

Readers are encouraged to review the Company’s Current Report on Form 8-K filed under Item 4.02 (Non-Reliance on Previously Issued Financial Statements or a Related Audit Report or Completed Interim Review), which describes the Company’s determination that its previously issued condensed consolidated financial statements for the quarter ended March 31, 2026 should no longer be relied upon and should be restated.

The information in this Item 2.02 and Exhibit 99.1 to this Current Report on Form 8-K shall not be deemed “filed” for purposes of Section 18 of the Securities Exchange Act of 1934, as amended, or otherwise subject to the liabilities of that section, nor shall it be deemed incorporated by reference in any filing under the Securities Act of 1933, as amended, except as expressly set forth by specific reference in such filing.

Item 7.01. Regulation FD Disclosure.

The information disclosed in Item 2.02 of this Current Report on Form 8-K, including Exhibit 99.1, is incorporated into this Item 7.01 by reference.

On August 13, 2026, the Company made available a presentation on its website. A copy of the presentation is attached hereto as Exhibit 99.2. Information contained on the Company’s website is not incorporated by reference into and should not be considered to be part of this Current Report on Form 8-K.

The information contained in this Item 7.01, including Exhibit 99.2 attached hereto, which is incorporated into this Item 7.01 by reference, is being furnished and shall not be deemed “filed” for 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 incorporated by reference in any filing under the Securities Act of 1933, as amended (the “Securities Act”), or the Exchange Act, except as expressly set forth by specific reference in such filing. The information set forth in this Item 7.01 and Exhibit 99.2 attached hereto shall not be deemed an admission as to the materiality of any information in this Current Report on Form 8-K that is required to be disclosed solely to satisfy the requirements of Regulation FD.

Forward-Looking Statements

Exhibit 99.2 attached hereto contains, and may indicate, forward-looking statements within the meaning of Section 27A of the Securities Act, Section 21E of the Exchange Act and as defined in the U.S. Private Securities Litigation Reform Act of 1995. Forward-looking statements include, but are not limited to, statements that express the Company’s intentions, beliefs, expectations, strategies, predictions or any other statements related to the Company’s future activities, or future events or conditions, including without limitation, those statements relating to the success of its products and product candidates, timing, progress and results of any preclinical and clinical trials, its estimates regarding the potential market opportunity for its products and product candidates, its ability to develop its pipeline, its ability to protect its intellectual property and enforce its intellectual property rights, its ability to procure new customers and partners, and its ability to execute its development strategy and sustain its competitive position. Actual future results and trends may differ materially depending on a variety of factors, including, but not limited to, the Company’s limited operating history, its ability to establish its market development capabilities to commercialize its products and generate any revenue, its ability to secure and execute financing transactions, and its ability to maintain regulatory approval of certain of its products, which can be identified by terminology such as “anticipate,” “believe,” “continue,” “could,” “estimate,” “expect,” “intend,” “may,” “plan,” “potential,” “predict,” “project,” “seek,” “should,” “target,” “will,” “would” and other similar expressions intended to identify forward-looking statements, although not all forward-looking statements contain these identifying words. These statements are not historical facts and are based on current expectations, estimates and projections about the Company’s business based, in part, on assumptions made by its management. These statements are not guarantees of future performance and involve risks, uncertainties and assumptions that are difficult to predict, many of which are beyond the Company’s control. Any forward-looking statements speak only as of the date on which they are made, and the Company undertakes no obligation to update any forward-looking statement to reflect events or circumstances after the date of this Form 8-K, except as required by applicable law.

Item 9.01. Financial Statements and Exhibits

(d) Exhibits:

Exhibit No.	Description
99.1	Press Release, dated August 13, 2026
99.2	Company 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: August 13, 2026

Pelthos Therapeutics Inc.

By: /s/ John M. Gay

Name: John M. Gay

Title: Chief Financial Officer



Pelthos Therapeutics Announces Second Quarter 2026 Financial Results

Zelsuvmi® net product revenue grew 45% to \$15.4 million during the second quarter of 2026 from \$10.7 million in the first quarter of 2026

11,925 Zelsuvmi units were dispensed by 4,571 unique prescribers during the second quarter of 2026, representing a 48% quarter-over-quarter increase in units dispensed

More than 25,000 patients have now been prescribed Zelsuvmi since it was commercially launched in July 2025

Management will host a conference call today, August 13, 2026, at 8:30 a.m. ET

DURHAM, N.C., August 13, 2026 — Pelthos Therapeutics Inc. (NYSE American: PTHS), a biopharmaceutical company committed to commercializing innovative therapeutic products for unmet patient needs (“Pelthos,” “we” or the “Company”), today announced its financial results for the second quarter ended June 30, 2026, which can be found in the Financial Results section of the Company’s website at <https://ir.pelthos.com/financial-info/financial-results>.

Second Quarter and Recent Highlights

- Zelsuvmi, the first at-home FDA-approved treatment (ages 1 and above) for molluscum contagiosum, a highly contagious viral skin infection that largely afflicts children, was launched in July 2025 and has generated \$42.3 million in net sales in the first four quarters of commercial operations.
- As of June 30, 2026, 29,126 units of Zelsuvmi have been dispensed and written by 7,414 unique prescribers. Units of Zelsuvmi dispensed increased from 8,084 in the first quarter of 2026 to 11,925 in the second quarter of 2026, representing a 48% increase. For the first full year, since its commercial launch in July of 2025 through the end of the second quarter of 2026, more than 25,000 patients have been prescribed Zelsuvmi.
- In January 2026, we entered into a \$50.0 million senior secured term loan facility with Horizon Technology Finance, of which we drew \$30.0 million at the close. Based on the Company achieving trailing twelve-month net product revenues of \$42.3 million as of June 30, 2026, we believe we have achieved access to an additional \$10.0 million under the Horizon facility, subject to the lender’s discretion.
- The Company’s cash balance as of June 30, 2026 was \$24.2 million, which based on its current projections, is expected to support the current business plan.
- As of June 30, 2026, we had approximately 9.0 million shares outstanding on an as-converted basis, which includes the conversion of approximately 52,128 shares of the Company’s Series A and 2,600 shares of Series C Convertible Preferred Stock, and approximately 3.7 million shares of common stock issued and outstanding.

Management Commentary

Scott Plesha, CEO of Pelthos, commented, “The growing adoption of Zelsuvmi and an increasing number of prescriptions written contributed significantly to our strong performance in the second quarter. We expect further growth for Zelsuvmi, with over 4,200 units dispensed in July and total units dispensed since the launch surpassing a significant milestone of 30,000. We believe that our continued commercial success, combined with the Horizon facility, and potential access to additional funds subject to the lender’s discretion, will provide us with the capital and flexibility necessary to advance our business plan. This includes the planned commercialization of Xepi® in the first quarter of 2027 and Xeglyze® in the third quarter of 2027.”

Second Quarter 2026 Financial Summary

- Net product revenue for Zelsuvmi during the second quarter of 2026 was \$15.4 million, as compared to \$10.7 million in the first quarter of 2026, representing a 45% quarter-over-quarter increase.
- Cost of goods sold was \$3.6 million for the second quarter of 2026 compared to \$1.7 million in the first quarter of 2026. Cost of goods sold includes fair value adjustments related to finished goods and active pharmaceutical ingredient inventory (“API”) on hand at the time of the Company’s merger in July 2025. Cost of goods sold for the second quarter of 2026 includes a \$0.9 million write-off of commercial API inventory identified through the Company’s routine in-process quality control and testing as narrowly falling outside specific tolerances for use in commercial drug product. The underlying procedural cause of these out-of-specification results was addressed and subsequent API manufacturing has commenced and met specifications.
- Selling, general and administrative (“SG&A”) expenses were \$27.7 million for the second quarter of 2026, as compared to \$21.1 million for the first quarter of 2026, representing a 31% quarter-over-quarter increase. Quarter-over-quarter changes in SG&A included: (i) an increase in royalty and milestone expense of \$6.0 million, due primarily to two non-recurring sales based milestones recorded during the second quarter in an amount of \$5.3 million, (ii) an increase in cash basis personnel costs of \$0.4 million, including \$0.5 million of non-recurring severance, (iii) an increase of \$0.7 million in non-cash expenses, comprised of stock based compensation and depreciation, and (iv) an increase of \$0.7 million in corporate expenses; offset by (v) a reduction in regulatory and manufacturing related expenses of \$0.8 million, and (vi) a reduction in marketing, travel, sales and commercial expenses of \$0.5 million.
- Interest expense for the second quarter of 2026 and first quarter of 2026, respectively, was \$2.4 million. Interest expense is attributable to (i) the Company’s existing convertible notes and its Horizon facility; and (ii) the accounting treatment of certain royalty and purchase agreement obligations entered into by the Company.
- Change in fair value of debt, related to the convertible notes issued in November 2025, was \$3.7 million in the second quarter of 2026, as compared to \$9.6 million for the first quarter of 2026. At issuance, the Company analyzed the terms of the convertible notes and its embedded features concluding it was appropriate to account for the convertible notes at fair value. Accordingly, the Company initially recognized the convertible notes at fair value and will subsequently measure the convertible notes at fair value with changes in fair value recorded in current period earnings or other comprehensive income.
- Net loss for the second quarter of 2026 was \$(23.4) million, as compared to \$(25.1) million for the first quarter of 2026.
- Adjusted EBITDA for the second quarter of 2026 was \$(5.7) million, as compared to \$(8.0) million for the first quarter of 2026, on a comparative basis discussed within the Non-GAAP Financial Information below.
- See additional detail within the Summary Financial Statement tables and Non-GAAP Financial Information below.

Webcast and Conference Call

Management will host a conference call and webcast today at 8:30 a.m. ET to discuss the Company’s second quarter 2026 results and provide a corporate update. Interested parties may participate in the call by dialing:

(877) 451-6152 (Domestic)
(201) 389-0879 (International)
Conference ID: 13761132

The live webcast will be accessible in the Investors section of the Company’s website or by following the direct link:

https://viaavid.webcasts.com/starthere.jsp?ei=1767131&tp_key=12eae87cb3

For those who cannot listen to the live broadcast, an online replay will be available in the Investors section of Pelthos' website.

About Pelthos Therapeutics

Pelthos Therapeutics is a commercial-stage biopharmaceutical company focused on building and advancing a portfolio of differentiated cutaneous infectious disease products that address unmet patient needs. Zelsuvmi® (berdazimer) topical gel, 10.3%, the company's lead product, is the first and only prescription therapy approved for use at home by patients, parents, and caregivers to treat molluscum contagiosum. The company's portfolio of assets includes Xepi® (ozenoxacin) Cream, 1%, a topical treatment for impetigo, and Xeglyze® (abametapir), a topical treatment for head lice. More information is available at www.pelthos.com. Follow Pelthos on [LinkedIn](#) and [X](#).

Forward-Looking Statements

This press release contains forward-looking statements, as defined in Section 21E of the Securities Exchange Act of 1934, regarding Pelthos' current expectations. All statements, other than statements of historical fact, could be deemed to be forward-looking statements. In some instances, words such as "plans," "believes," "expects," "anticipates," and "will," and similar expressions, are intended to identify forward-looking statements. Readers are cautioned not to place undue reliance on these forward-looking statements, which reflect our good faith beliefs (or those of the indicated third parties) and speak only as of the date hereof. These forward-looking statements include, without limitation, references to our expectations regarding (i) our belief that we have achieved access to an additional \$10.0 million under the Horizon facility and the likelihood that the lender, in its discretion, will grant access to such additional \$10.0 million; (ii) the anticipated ongoing growth for Zelsuvmi; (iii) our belief that our continued commercial performance, paired with the Horizon facility and potential access to additional funds, subject to the lender's discretion will provide us with the capital and flexibility needed to advance our business plan; (iv) the potential timing for the commercialization and anticipated launch of Xepi and Xeglyze; (v) our belief that the exclusion of certain items in calculating Adjusted EBITDA can provide a useful measure for period-to-period comparisons of our business; and (vi) our belief that Adjusted EBITDA provides useful information to investors in understanding and evaluating our operating results. These statements are not guarantees of future performance and are subject to certain risks, uncertainties and assumptions that are difficult to predict. Factors that could cause actual results to differ materially from those set forth in such forward-looking statements include, but are not limited to, risks and uncertainties related to there being no guarantee that the trading price of the combined company's Common Stock will be indicative of the combined company's value or that the combined company's Common Stock will become an attractive investment in the future; we may rely on collaborative partners for milestone payments, royalties, materials revenue, contract payments and other revenue projections and may not receive expected revenue; we and our partners may not be able to timely or successfully advance any product(s) in our internal or partnered pipeline or receive regulatory approval and there may not be a market for the product(s) even if successfully developed and approved; and changes in general economic conditions, including as a result of war, conflict, epidemic diseases, the implementation of tariffs, and ongoing or future litigation could expose us to significant liabilities and have a material adverse effect on us. These and other risks and uncertainties are described more fully in our filings with the U.S. Securities and Exchange Commission. The information in this press release is provided only as of the date of this press release, and we undertake no obligation to update any forward-looking statements contained in this press release based on new information, future events, or otherwise, except as required by law.

Contacts

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Summary Financial Statements
Pelthos Therapeutics Inc.
Selected Condensed Consolidated Balance Sheet Data
(unaudited)
(in thousands)

	June 30, 2026	December 31, 2025
Cash and cash equivalents	\$ 24,192	\$ 17,973
Accounts receivable, net	14,482	8,858
Inventory	21,472	23,574
Total current assets	63,356	53,410
Total assets	137,356	130,397
Accounts payable	\$ 5,853	\$ 2,986
Accrued expenses	20,745	15,364
Total current liabilities	31,959	25,993
Total liabilities	138,857	91,516
Total stockholders' (deficit) equity	\$ (1,501)	\$ 38,881
Total liabilities and stockholders' equity	137,356	130,397

Pelthos Therapeutics Inc.
Condensed Consolidated Statements of Operations
(unaudited)
(in thousands except share and per share data)

	Three Months Ended June 30,		Six Months Ended June 30,	
	2026	2025	2026	2025
Revenue				
Net product revenues	\$ 15,420	\$ —	\$ 26,085	\$ —
License and collaboration revenues	183	—	424	—
Total revenue	15,603	—	26,509	—
Operating expenses				
Cost of goods sold	3,610	—	5,283	—
Selling, general and administrative	27,673	2,716	48,777	4,356
Research and development	600	515	786	709
Amortization of intangible assets	1,043	—	2,074	—
Total operating expenses	32,926	3,231	56,920	5,065
Operating loss	(17,323)	(3,231)	(30,411)	(5,065)
Other (expense) income				
Interest expense	(2,389)	(218)	(4,742)	(352)
Change in fair value of convertible debt	(3,705)	—	(13,337)	—
Total other expense	(6,094)	(218)	(18,079)	(352)
Net loss before provision for income taxes	(23,417)	(3,449)	(48,490)	(5,417)
Provision for income taxes	—	—	—	—
Net loss	\$ (23,417)	\$ (3,449)	\$ (48,490)	\$ (5,417)
Net loss per common share - basic and diluted	\$ (6.62)	\$ (5.29)	\$ (14.15)	\$ (8.56)
Weighted average number of common shares outstanding - basic and diluted	3,539,464	651,921	3,426,232	632,513

The table below sets forth the income statement for the second quarter of 2026 and the first quarter of 2026.

Pelthos Therapeutics Inc.
Condensed Consolidated Statements of Operations
(unaudited)
(in thousands except share and per share data)

	Three Months Ended	
	June 30, 2026	March 31, 2026 (As Restated)
Revenue		
Net product revenues	\$ 15,420	\$ 10,665
License and collaboration revenues	183	241
Total revenue	15,603	10,906
Operating expenses		
Cost of goods sold	3,610	1,673
Selling, general and administrative	27,673	21,104
Research and development	600	186
Amortization of intangible assets	1,043	1,031
Total operating expenses	32,926	23,994
Operating loss	(17,323)	(13,088)
Other (expense) income		
Interest expense	(2,389)	(2,353)
Change in fair value of convertible debt	(3,705)	(9,632)
Total other expense	(6,094)	(11,985)
Net loss before provision for income taxes	(23,417)	(25,073)
Provision for income taxes	—	—
Net loss	\$ (23,417)	\$ (25,073)
Net loss per common share - basic and diluted	\$ (6.62)	\$ (7.57)
Weighted average number of common shares outstanding - basic and diluted	3,539,464	3,311,742

Non-GAAP Financial Information

Adjusted EBITDA

To provide investors with additional information regarding the Company's financial results, we have provided within this press release Adjusted EBITDA, a non-GAAP financial measure. We define Adjusted EBITDA as net loss adjusted to eliminate (i) stock-based compensation expense, (ii) the inventory valuation step-up recognized in cost of goods sold resulting from the July 1, 2025 acquisition of LNHC, Inc., as described below, (iii) change in fair value of convertible debt, (iv) non-recurring milestones due to license agreement counterparties, (v) interest expense, (vi) amortization of intangible assets, (vii) depreciation expense, and (viii) the provision for income taxes. We have provided a reconciliation below of Net Loss, the most directly comparable GAAP financial measure, to Adjusted EBITDA.

The Company accounts for business acquisitions using the acquisition method of accounting in accordance with Accounting Standards Codification ("ASC") 805, Business Combinations. ASC 805 requires, among other things, that assets acquired and liabilities assumed be recognized at their fair values, as determined in accordance with ASC 820, Fair Value Measurements ("ASC 820"), as of the acquisition date. As part of the July 1, 2025 acquisition of LNHC, Inc., the fair value of the inventory acquired was estimated using the top/down method that considers the estimated selling price, costs to complete, disposal costs, profit margin on disposal effort, and holding costs. Significant assumptions include management's estimates for the selling price and the costs to be incurred related to

the disposal effort of the inventory. The non-cash inventory valuation step-up from the acquisition of LNHC, Inc. is recognized within cost of goods sold in the periods presented.

We have included Adjusted EBITDA in this press release because it is a key measure used by our management to understand and evaluate our core operating performance and trends, to prepare and approve our annual budget and to develop short- and long-term operating plans. In particular, we believe the exclusion of certain items from net loss in calculating Adjusted EBITDA can provide a useful measure for period-to-period comparisons of our business. Accordingly, we believe that Adjusted EBITDA provides useful information to investors in understanding and evaluating our operating results. Our use of Adjusted EBITDA has limitations as an analytical tool, and you should not consider it in isolation or as a substitute for analysis of our results as reported under GAAP

The following table presents a reconciliation of Net Loss to Adjusted EBITDA for each of the periods indicated (in thousands):

	Three Months Ended June 30,		Six Months Ended June 30,	
	2026	2025	2026	2025
Net loss	\$ (23,417)	\$ (3,449)	\$ (48,490)	\$ (5,417)
Adjustments:				
Stock-based compensation	2,760	394	4,668	850
Cost of goods sold basis step-up	2,295	—	3,934	—
Change in fair value of convertible debt	3,705	—	13,337	—
Non-recurring milestones	5,250	—	5,250	—
Interest expense	2,389	218	4,742	352
Amortization of intangible assets	1,043	—	2,074	—
Depreciation	278	—	747	—
Adjusted EBITDA	<u>\$ (5,697)</u>	<u>\$ (2,837)</u>	<u>\$ (13,738)</u>	<u>\$ (4,215)</u>



Corporate Presentation

AUGUST 2026

Legal Disclaimer

This presentation of Pelthos Therapeutics Inc. ("we", "us", "our" or the "Company") contains "forward-looking statements" within the meaning of the Private Securities Litigation Reform Act and other securities laws. Words such as "anticipate," "believe," "continue," "could," "estimate," "expect," "intend," "may," "plan," "potential," "predict," "project," "seek," "should," "target," "will," "would" or similar expressions and the negatives of those term are intended to identify forward-looking statements. Forward-looking statements reflect management's current expectations, are based on judgments and assumptions, are inherently uncertain and are subject to risks, uncertainties and other factors, which could cause the Company's actual results, performance or achievements to differ materially from expected future results, performance or achievements expressed or implied in those forward-looking statements. Examples of these forward-looking statements and the related risks, uncertainties and other factors include, but are not limited to, the following: the success of the launch of the products in our portfolio, timing, progress and results of any preclinical and clinical trials, our estimates regarding the potential market opportunity for our products, our ability to develop our pipeline, our ability to protect our intellectual property and enforce our intellectual property rights, and our ability to execute our development strategy and sustain our competitive position. Actual future results and trends may differ materially depending on a variety of factors, including, but not limited to, the Company's limited operating history, the Company's ability to establish our market development capabilities to commercialize our products and generate any revenue, and the Company's ability to maintain regulatory approvals of products in our portfolio.

Forward-looking statements are provided to allow potential investors the opportunity to understand management's beliefs and opinions in respect of the future so that they may use such beliefs and opinions as one factor in evaluating an investment. These statements are not guarantees of future performance and undue reliance should not be placed on them. Any forward-looking statement in this presentation, in any related presentation supplement and in any related free writing presentation reflects our current view with respect to future events and is subject to these and other risks, uncertainties and assumptions relating to our business, results of operations, industry and future growth. You should read this presentation with the understanding that our actual future results may be materially different from any future results expressed or implied by these forward-looking statements. Except as required by law, we assume no obligation to update or revise these forward-looking statements for any reason, even if new information becomes available in the future.

Investment Highlight

- ✓ Commercial biopharmaceutical company focused on growing, differentiated cutaneous infections product portfolio
- ✓ Highly synergistic Xepi and Xeglyze product acquisitions leverage Zelsuvmi's current commercial and market access team and infrastructure
- ✓ Strong potential revenue streams with >20,000 Zelsuvmi units dispensed from commercial launch in July 2025 through April 2026
- ✓ Experienced management team to manage execution

Product Portfolio



- Large addressable market with \$2,008.50 wholesale acquisition cost ("WAC")
- **Launched July 2025**
- Modest acquisition cost, unencumbered future revenue stream
- **Expected launch Q1 2027**
- 6–12 million U.S. cases annually
- **Expected launch Q3 2027**

Corporate Profile

Pelthos is a competitive drug portfolio company — committed to commercializing innovative, safe, and efficacious therapeutic products to help patients with unmet dermatological treatment burdens

Zelsuvmi: Launched in July 2025.

- First and only at home treatment addressing *Molluscum contagiosum* ("MC"), a large, underserved market treating contagious viral disease

Recent portfolio acquisitions: Two FDA-approved complimentary dermatological acquisitions, will leverage Zelsuvmi commercial infrastructure buildout

- **Xepi** (ozenoxacin) Cream 1% - novel topical treatment for impetigo
 - First line impetigo treatment addresses antimicrobial resistance in pediatric dermatology, drug relaunch expected Q1 2027
- **Xeglyze** (abametapir) Lotion 0.74% - novel topical treatment for head lice
 - Commercial launch expected Q3 2027

Experienced management team: Over 20 successful prior drug launches, including Cosentyx, Otezla, Ohtuvayre, Xifaxan

Key Data Points (as of 08/10/26, except where noted)

Ticker	PTHS
Stock Price	\$29.00
O/S Shares of Common Stock (with Pref A conversion)	~9.0M
Market Cap	~\$260M
60-Day Avg. Daily Trading Volume	~51,000 shares (Yahoo)
Cash at end of Q2 2026	\$24.2M

Management Team



Scott Plesha | Chief Executive Officer

- >30 years of experience in the pharmaceutical industry, including two decades building and leading specialty pharma commercial efforts
- President and Chief Commercial Officer at BDSI until it was acquired by Collegium Pharmaceutical in 2022
- Grew BDSI sales from \$5 million to \$160 million
- Previously served as Senior Vice President of Gastrointestinal Sales at Salix Pharmaceuticals. During fifteen-year tenure at Salix, led nationwide salesforce that grew product sales to more than \$1.5 billion annually
- Earned a BA in Pre-Medicine and Pre-Medical Studies at DePauw University and pursued graduate studies in Dentistry at Indiana University Dental School



John M. Gay | Chief Financial Officer

- >25 years of experience in public company finance and accounting for life science and technology companies, including R&D and commercial organizations
- Raised nearly \$300 million in capital and managed various acquisitions, divestitures, integrations and initial public offerings
- Corporate Controller for Furiex Pharmaceuticals, Inc., from its initial spin-out transaction, prior to the sale to Forest Labs for \$1.1 billion
- Earned BAs in Economics and History, and a Master's in Accounting from the University of North Carolina at Chapel Hill



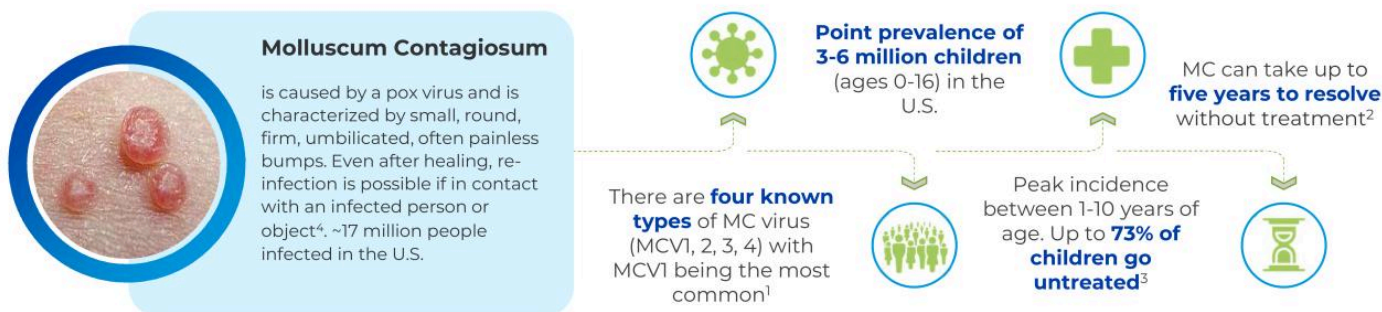
Sai Rangarao | Chief Commercial Officer

- >18 years of experience leading, launching, and marketing pharmaceutical products
- VP of Marketing & Head of Neurology Sales at Collegium Pharmaceutical
- VP of Marketing & Commercial Operations at BDSI, until it was acquired by Collegium in 2022
- Head of US Dermatology Marketing for Otezla at Celgene leading to acquisition by Amgen for \$13 Billion
- Member of the commercial and marketing organization at Novartis Pharmaceuticals that launched COSENTYX® in the U.S
- Earned an MS in Bioscience Regulatory Affairs from The Johns Hopkins University, an MBA and MS from the New Jersey Institute of Technology, and a BS in Computer Science from Indiana University of Pennsylvania

Molluscum & Zelsuvmi Overview

Molluscum Contagiosum

A highly infectious viral condition primarily affecting children 1 year of age or older



Untreated Molluscum Contagiosum Has Severe Effects

Infection, Persistence, and Spread

Auto-inoculation²

Highly contagious to others

+ risk of secondary bacterial infections²

Potential worsening of atopic dermatitis

Itching, redness

Inflammation

Anxiety

Social withdrawal

Pain & Skin Irritation

Visible and Psychological Impacts

1) Hebert AA, Bhatia N, Del Rosso JQ. Molluscum Contagiosum: Epidemiology, Considerations, Treatment Options, and Therapeutic Gaps. J Clin Aesthet Dermatol. 2023 Aug;16(8 Suppl 2):S4-S11. PMID: 37636018; PMCID: PMC10453394. 2) Ludmann P. American Academy of Dermatology. Molluscum contagiosum. 4 October 2023. 3) Basdag H, Rainer BM, Cohen BA. Molluscum contagiosum: to treat or not to treat? Experience with 170 children in an outpatient clinic setting in the northeastern United States. Pediatr Dermatol. 2015;32(3):353-357. doi:10.1111/pde.12504. 4) Schaffer JV, Berger EM. Molluscum Contagiosum. JAMA Dermatol. 2016;152(9):1072. doi:10.1001/jamadermatol.2016.2367. 5) CDC. Clinical Overview of Molluscum Contagiosum. Jan 2025

Zelsuvmi Has the Potential to Shift MC Treatment Paradigm

The 1st & Only At Home Prescription Treatment

Previous Treatment Options



- Other available topical treatment **requires in-office visits every 3 weeks**²



- **Painful, destructive** treatments³



- Necessitates travel to HCP offices, adding to the **time burden for MC patients and caregivers**²



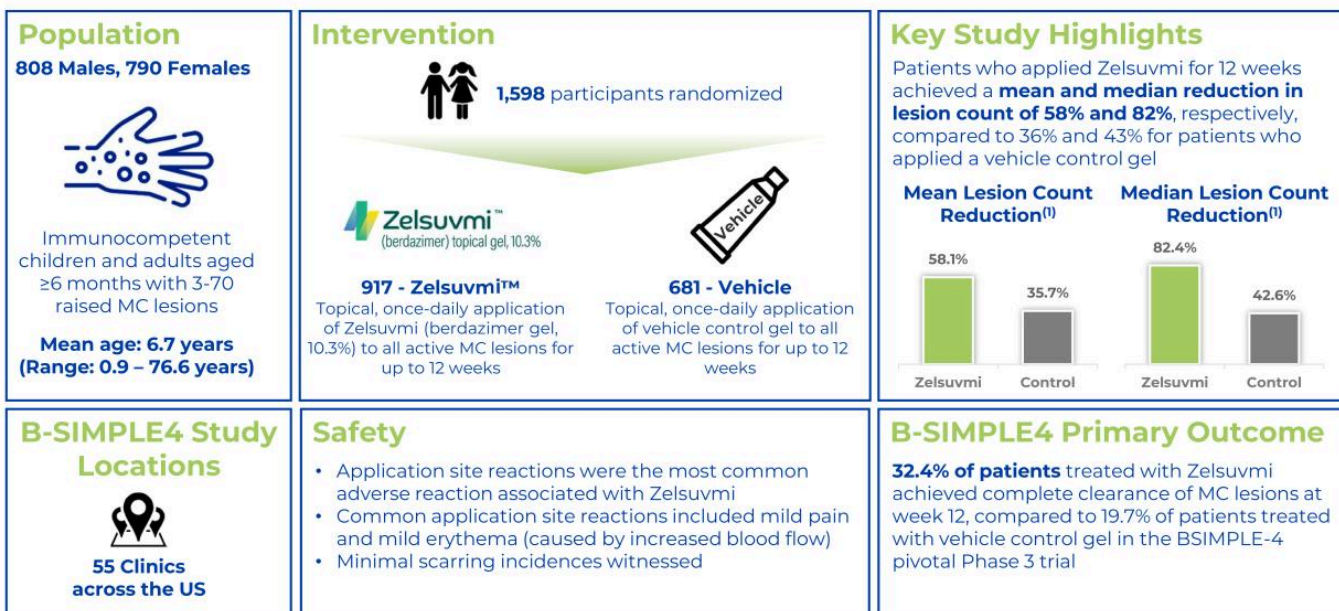
- Remaining treatment options such as off-label drugs / natural remedies have **unproven efficacy**⁴



- **Daily** application that can be **started immediately**
- **Attractive safety profile** demonstrated in clinical trials with no / minimal scarring^{5,6}
- **First FDA approved medication** for molluscum that can be applied at home by patients or caregivers⁵
- **Demonstrated, proven efficacy** across key primary and secondary endpoints in clinical trials⁶

1) Least-squares mean count reduction. See Figure 9: Browning JC, Hebert A, Enloe C, Cartwright M, Maeda-Chubachi T. Berdazimer Gel 10.3% is a Clinically Meaningful Therapeutic Intervention for Molluscum Contagiosum. Abstract and poster presented at Fall Clinical 2024, Las Vegas, NV, October 26-27, 2024. 2) Eichenfield LF, Kwong P, Gonzalez ME, et al. Safety and Efficacy of VP-102 (Cantharidin, 0.7% w/v) in Molluscum Contagiosum by Body Region: Post hoc Pooled Analyses from Two Phase III Randomized Trials. J Clin Aesthet Dermatol. 2023;14(10):42-47. 3) Hebert AA, Bhatia N, Del Rosso JQ. Molluscum Contagiosum: Epidemiology, Considerations, Treatment Options, and Therapeutic Gaps. J Clin Aesthet Dermatol. 2023;16(8 Suppl 1):S4-S11. 4) Ong SK, Hoft I, Siegfried E. Analysis of over-the-counter products marketed to treat molluscum contagiosum. Pediatr Dermatol. 2021;38(5):1400-1403. doi:10.1111/pde.14776. 5) Zelsuvmi Package Insert. 6) Sugarman JL, Hebert A, Browning JC, et al. Berdazimer gel for molluscum contagiosum: An integrated analysis of 3 randomized controlled trials. J Am Acad Dermatol. 2024;90(2):299-308. doi:10.1016/j.jaad.2023.09.066Ong

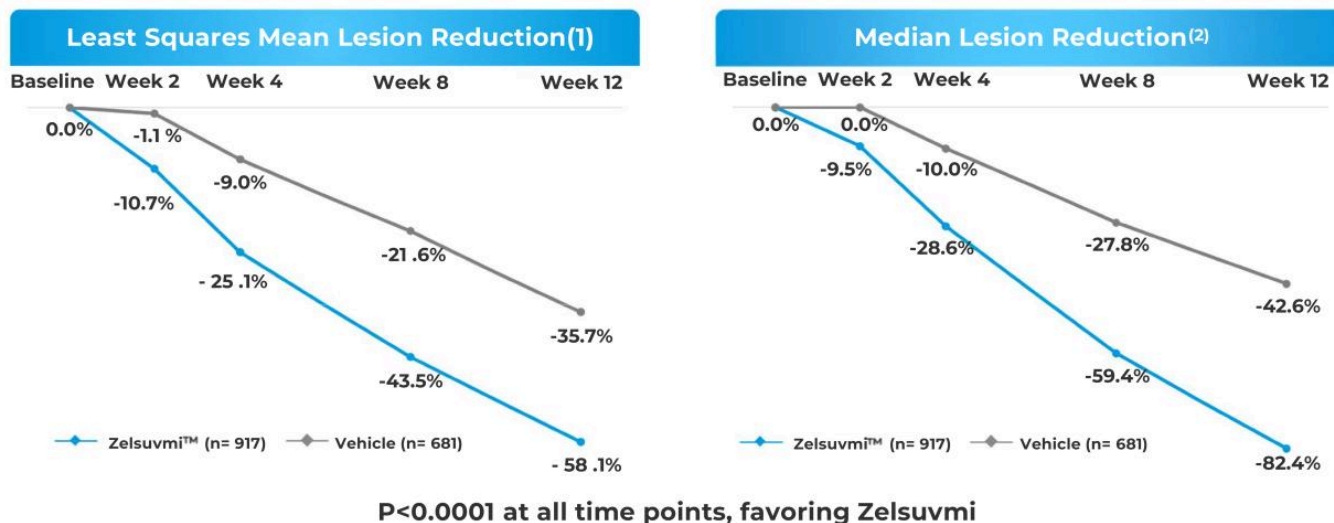
Zelsuvmi Efficacy Shown in Phase 3 Clinical Trials Drives Commercial Launch



⁽¹⁾ p-value <0.0001, favoring Zelsuvmi™.
Source: Sugarman JL, Hebert A, Browning JC, Paller AS, Stripling S, Green LJ, Cartwright M, Enloe C, Wells N, Maeda-Chubachi T. Berdazimer gel for molluscum contagiosum: An integrated analysis of 3 randomized controlled trials. J Am Acad Dermatol. 2023 Oct 5;S0190-9622(23)02890-6. doi:10.1016/j.jaad.2023.09.066.Epub ahead of print. PMID: 37804936.

Phase 3 Trial Results

Zelsuvmi showed statistically significant benefit vs. vehicle after 2 weeks of therapy and through out the entire 12-week length of the Phase 3 studies



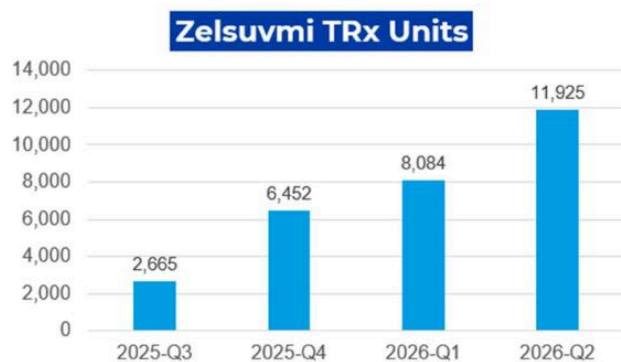
1) Figure 9: Browning JC, Hebert A, Enloe C, Cartwright M, Maeda-Chubachi T. Berdazimer Gel 10.3% is a Clinically Meaningful Therapeutic Intervention for Molluscum Contagiosum. Abstract and poster presented at Fall Clinical 2024. Las Vegas, NV. October 24-27, 2024. 2) Figure 10: Browning JC, Hebert A, Enloe C, Cartwright M, Maeda-Chubachi T. Berdazimer Gel 10.3% is a Clinically Meaningful Therapeutic Intervention for Molluscum Contagiosum. Abstract and poster presented at Fall Clinical 2024. Las Vegas, NV. October 24-27, 2024.

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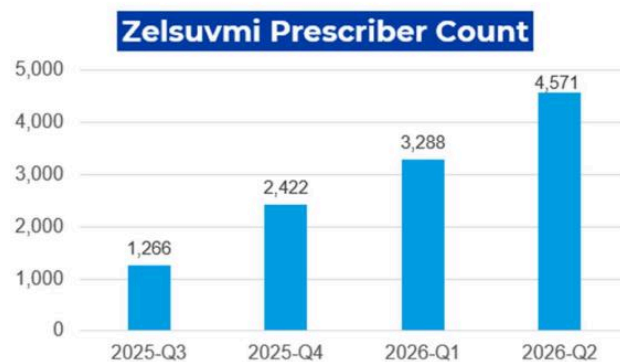
Zelsuvmi Commercial Overview

Strong Zelsuvmi Quarter over Quarter Growth Continues

Robust growth into first full year of launch



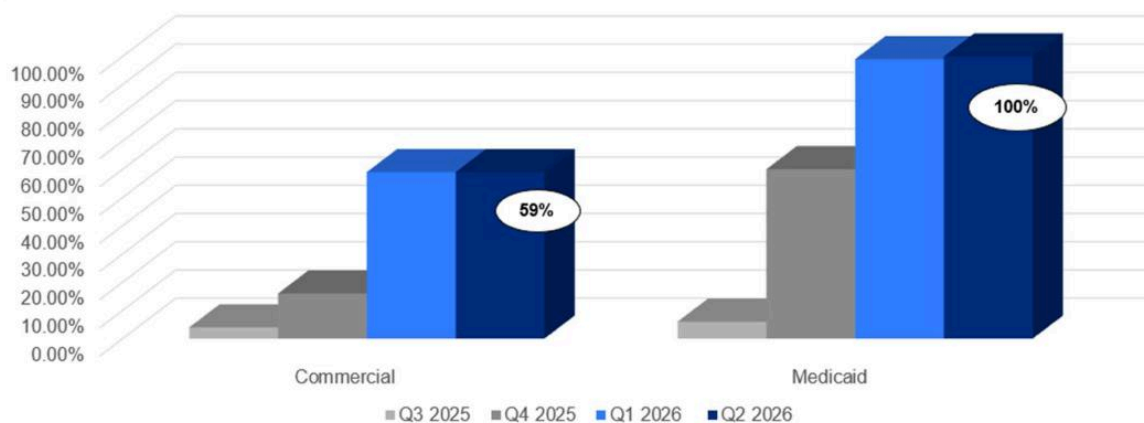
Total of 29,126 Prescribed Units



Total of 7,414 Unique Prescribers

Data Source: Symphony Health- Metys Data

Zelsuvmi Covered Lives by Quarter*

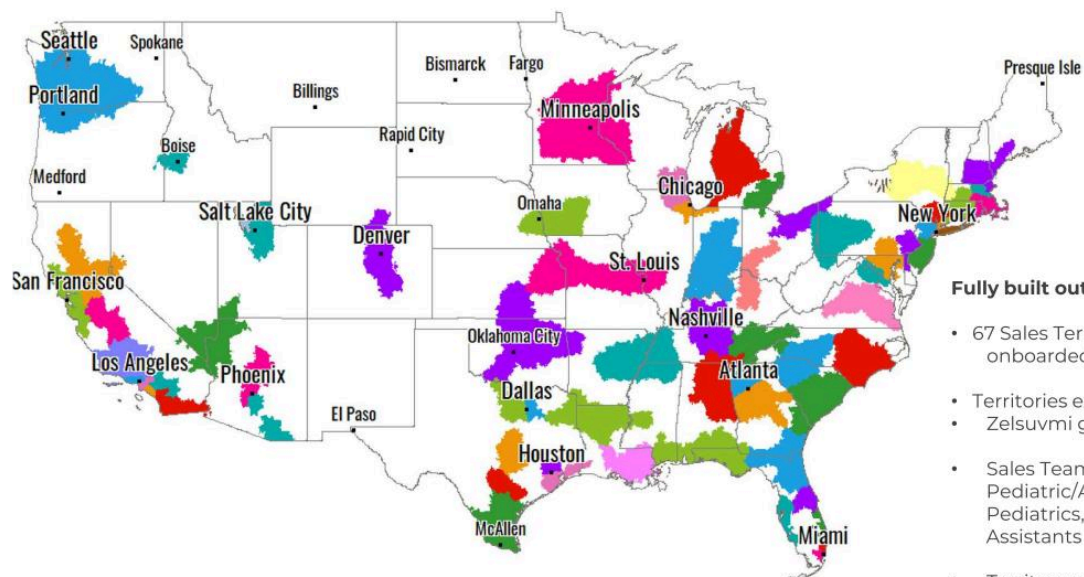


- Selective contracting strategy
- 70% combined Medicaid/commercial coverage with 1 contract
- Favorable Gross to Nets
- Favorable approval rates with all payers

- Commercial, Cash and Assistance Programs - 72% of TRxs
- Managed and FSS Medicaid - 28%
- Medicare <1% of TRxs

*Clarivate (DRC) Fingertip Formulary

Strong Sales Team Execution



Fully built out commercial team:

- 67 Sales Territories Expansion completed & onboarded covering ~53% of MC Claims
- Territories expansion complete, contributing to Zelsuvmi growth
- Sales Team targeting:
Pediatric/Adult Dermatologists, General Pediatrics, Nurse Practitioners & Physicians Assistants
- Territory managers supported by Sales Training, Marketing, Commercial Operations & Market Access teams

Comprehensive Zelsuvmi Tactical Execution



National & Regional Conference Presence



YouTube Promotional Commercial



New Patient Testimonials & HCP Video Series



Digital Marketing



Live & Virtual Educational Speaker Development

Zelsuvmi
GO

ZELSUVMI GO Patient Support Program

Xepi: New Product Acquisition



Xepi (ozenoxacin) Cream for the treatment of Impetigo



Acquired from BioFrontera
in October 2025

FDA Approved in 2017

Exclusivity until **2032**

Xepi Clinical Story

- Ozenoxacin cream 1% developed as first line treatment in patients aged 2 months and older
- 15 clinical studies in Phase 1 & 2 conducted
- Two Pivotal Phase 3 studies conducted in both adult & pediatric patients with impetigo 2 months old and up
- Ozenoxacin demonstrated superior clinical and bacteriological outcomes vs. vehicle control

Impetigo Facts¹

- #1 bacterial infection seen in pediatrician offices, represents 1-2% of all visits to Pediatricians in the US, with **135M** children suffering worldwide
- Impetigo is a highly contagious bacterial skin infection, most often caused by *Staphylococcus aureus* and/or Group A *Streptococcus* (*Streptococcus pyogenes*)
- Mupirocin resistance is growing significantly in the US

Pelthos Opportunity

- Strong synergy with existing commercial infrastructure for Zelsuvmi
- Significant overlap between Xepi & Zelsuvmi HCP call points
- Promotional alignment across Sales, Marketing & Commercial Operations
- Anticipated Commercial Launch: Q1 2027

Sources: CDC Website, Xepi Pack Insert, FDA.gov

¹<https://www.cdc.gov/group-a-strep/about/impetigo.html>

Xeglyze: New Product Acquisition



Xeglyze (abametapir) Lotion for the Treatment of Head Lice



Acquired from Hatchtech in
December 2025

FDA Approved in 2020

Exclusivity until **2034**

Xeglyze Clinical Story

- Abametapir lotion 0.74% developed as first line treatment in patients aged 6 months of age and older
- Phase 2b study completed in 2014 demonstrated 100% ovicidal efficacy
- Two Pivotal Phase 3 studies demonstrated that a single, ten-minute application of Xeglyze® results in a statistically significant increase in the proportion of subjects who are cleared of lice versus vehicle.

Head Lice Facts¹

- **100m+** infestations globally, with **6-12m cases** in the US, each year with substantial social cost
- Increasing resistance to current products containing pyrethrin, permethrin & malathion
- Existing products are only effective against lice and not eggs, and most require repeat treatments to break life cycle of infestation, leading to poor compliance and reduced efficacy

Pelthos Opportunity

- Strong synergy with existing commercial infrastructure for Zelsuvmi and Xepi
- Significant overlap between Xeglyze & Zelsuvmi HCP call points
- Promotional alignment across Sales, Marketing & Commercial Operations
- Anticipated Commercial Launch: Q3 2027

Sources: CDC Website, Xepi Pack Insert, FDA.gov

¹https://www.accessdata.fda.gov/drugsatfda_docs/label/2017/2089451bl.pdf

Summary Financial Statements

Summary Balance Sheet

	June 30, 2026	December 31, 2025
Cash and cash equivalents	\$ 24,192	\$ 17,973
Accounts receivable, net	14,482	8,858
Inventory	21,472	23,574
Total current assets	63,356	53,410
Total assets	137,356	130,397
Accounts payable	\$ 5,853	\$ 2,986
Accrued expenses	20,745	15,364
Total current liabilities	31,959	25,993
Total liabilities	138,857	91,516
Total stockholders' (deficit) equity	\$ (1,501)	\$ 38,881
Total liabilities and stockholders' equity	137,356	130,397

Notes

- Cash balance included \$30 million raised with Horizon Loan Facility in January 2026 with issuance of 5-year term notes; potential access to \$10 million as of June 30, 2026, subject to lender discretion; additional tranches totaling \$10 million available on hitting certain additional milestones
- Working capital of \$31.4 million as of June 30, 2026
- Strong stockholder equity position and cash balance provide resources to execute on our business plan

Summary Income Statement

	Three Months Ended	
	June 30, 2026	March 31, 2026 (As Restated)
Revenue		
Net product revenues	\$ 15,420	\$ 10,665
License and collaboration revenues	183	241
Total revenue	15,603	10,906
Operating expenses		
Cost of goods sold	3,610	1,673
Selling, general and administrative	27,673	21,104
Research and development	600	186
Amortization of intangible assets	1,043	1,031
Total operating expenses	32,926	23,994
Operating loss	(17,323)	(13,088)
Other (expense) income		
Interest expense	(2,389)	(2,353)
Change in fair value of convertible debt	(3,705)	(9,632)
Total other expense	(6,094)	(11,985)
Net loss before provision for income taxes	(23,417)	(25,073)
Provision for income taxes	—	—
Net loss	\$ (23,417)	\$ (25,073)

Notes

- Net product revenue increased 45% quarter over quarter
- SG&A expenses increased 31% quarter over quarter, related primarily to one-time milestones of ~\$5.3 million and royalty increase of ~\$0.8 million
- SG&A includes milestones & royalties and non-cash expenses (stock-based compensation and depreciation)
- SG&A spend supporting current and future net revenue growth

Appendix

Nitricil Platform Pipeline*

Indication	Asset Description	Approximate Time to NDA Filing
SB414 (AD/Psoriasis)	Berdazimer topical cream, dose TBD, for treatment of mild to moderate atopic dermatitis. Phase 1/2 Clinical stage.	7.5 years
SB208 (Tinea Pedis -> Onychomycosis)	Low alcohol berdazimer topical gel for treatment of athlete's foot with label expansion for onychomycosis following initial approval. Phase 2/3 Clinical stage.	5 years (T. Pedis) 6.5 years (Onychomycosis)
SB208 (Tinea Pedis + Onychomycosis)	Low alcohol berdazimer topical gel for treatment of both athlete's foot and onychomycosis. Phase 2/3 Clinical stage.	6.5 years
SB207 (EGW/PAW)	Berdazimer topical gel, 10.3% for treatment of external genital and perianal warts. Same active gel (Tube A) as Zelsuvmi but different hydrogel (Tube B) formulation. Phase 3 clinical stage.	6.5 years

*Pelthos has contractual rights to SB207 and would need to enter into a separate license for other indications set forth herein

NaV1.7 Pipeline

Indication	Asset Description	Approximate Time to NDA Filing
CT2000 Eye Drops Chronic Ocular Pain	CC8464 1%, 1.25% and 1.5% ophthalmic solution Phase 1-2a ready	3-4 years
CT2000 Eye Drops Acute Ocular Pain	CC8464 1%, 1.25% and 1.5% ophthalmic solution Phase 1-2a ready	2-3 years
CT3000 depot Nerve Blocks	CC8464 5% and 10% depot injectable Preclinical Stage	5+ years
CC8464 Oral Erythromelalgia	CC8464 melt-granulation capsules 50mg, 100mg, 400mg Phase 2 Ready	5+ years
CC8464 Oral Small Fibre Neuropathy	CC8464 melt-granulation capsules 50mg, 100mg, 400mg Phase 2 Ready	5+ years
CC8464 Oral Acute Pain	CC8464 melt-granulation capsules 50mg, 100mg, 400mg Phase 2 Ready	5+ years

Board of Directors



Peter Greenleaf, Chairman



Richard Baxter



Todd Davis



Andrew Einhorn



Ezra Friedberg



Richard Malamut, MD



Matt Pauls



Scott Plesha



Key Highlights

	Portfolio of FDA Approved Products	Differentiated portfolio of novel, cutaneous infectious disease products, including Zelsuvmi, Xepi and Xeglyze for the treatment of MC, impetigo and head lice, respectively
	Significant Unmet Need and Large Market Opportunities	Each Pelthos product is differentiated from existing treatment options with considerable market opportunities
	Barriers to Entry	Strong patent portfolio, along with complex, proprietary manufacturing process for Zelsuvmi and complex, multi-step manufacturing process for Xepi provides hefty market protection
	Operating Leverage	All three products utilize the same sales team, with largely overlapping call points, provides greater operating and financial leverage with very little dedicated overhead
	Strong Financial Position	Current balance sheet, revenue growth and strong existing investor support with substantial investable cash provides robust foundation for growth
	Biopharmaceutical Platform Poised for Growth	Strategically positioned to explore and integrate synergistic acquisitions, serving as a platform for investors seeking a strong foothold in the specialty biopharmaceutical market
	Pipeline	Opportunity to exploit legacy Channel clinical programs and work with Ligand to execute on clinical stage programs based on the same Nitricil platform as Zelsuvmi



Thank You



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