#### 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): September 2, 2025

#### PELTHOS THERAPEUTICS INC.

(Exact name of registrant as specified in its charter)

Nevada	001-41964	86-3335449
(State or other jurisdiction of incorporation)	(Commission File Numb	per) (IRS Employer Identification No.)
4020 Stirrup	p Creek Drive, Suite 110	
Ĭ	Durham, NC	27703
(Address of registra	ant's principal executive office)	(Zip code)
Registrant's	telephone number, including area code: (9	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 intend General Instruction A.2. below):	led to simultaneously satisfy the filing ob	ligation of the registrant under any of the following provisions (see
$\hfill \Box$ Written communications pursuant to Rule 425 under the Secur	ities Act (17 CFR 230.425)	
$\hfill \square$ Soliciting material pursuant to Rule 14a-12 under the Exchang	e Act (17 CFR 240.14a-12)	
☐ Pre-commencement communications pursuant to Rule 14d-2(b	under the Exchange Act (17 CFR 240.14	4d-2(b))
☐ Pre-commencement communications pursuant to Rule 13e-4(c	) under the Exchange Act (17 CFR 240.13	e-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 grow the Securities Exchange Act of 1934 (§240.12b-2 of this chapter).	wth company as defined in Rule 405 of th	e Securities Act of 1933 (§230.405 of this chapter) or Rule 12b-2 of
Emerging growth company $\boxtimes$		
If an emerging growth company, indicate by check mark if the reaccounting standards provided pursuant to Section 13(a) of the Ex		ed transition period for complying with any new or revised financial

#### Item 7.01. Regulation FD Disclosure.

On September 2, 2025, Pelthos Therapeutics Inc. (the "Company") issued a press release announcing that its Chief Executive Officer, Scott Plesha, and Chief Financial Officer, Francis Knuettel II, will present at the Wells Fargo 2025 Healthcare Conference in Boston, Massachusetts. Management's presentation will take place on Wednesday, September 3, 2025 at 1:30 p.m. Eastern Time. A webcast of the presentation will be available on the Events and Presentations pages of the Company's website at https://pelthos.com. Archived replays will be available for 90 days following the conference. A copy of the press release is furnished as Exhibit 99.1 to this Current Report on Form 8-K.

At the conference, the Company will provide an update on its product pipeline and discuss recent business developments, including its commercial launch of ZELSUVMI<sup>TM</sup>. The presentation materials furnished herewith as Exhibit 99.2 (the "Presentation Materials") will be used in connection with such presentation, are incorporated into this Item 7.01 by reference and will be posted on the Company's website at https://pelthos.com. 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 Exhibits 99.1 and 99.2 attached hereto, which are 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 Exhibits 99.1 and 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 forward-looking statements, as defined in Section 21E of the Securities Exchange Act of 1934, regarding the Company's 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) the success of the launch for ZELSUVMI, (ii) the Company's ability to develop its pipeline, (iii) the Company's ability to protect its intellectual property and to enforce its intellectual property rights, and (iv) the Company's ability to execute its development strategy and sustain its competitive position. 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 Current Report on Form 8-K is provided only as of the date of this Current Report on Form 8-K, and we undertake no obligation to update any forward-looking statements contained in this Current Report on Form 8-K based on new information, future events, or otherwise, except as required by law

#### Item 9.01. Financial Statements and Exhibits

(d) Exhibits:

Exhibit No.	Description
<u>99.1</u>	Press Release, dated September 2, 2025.
<u>99.2</u>	Pelthos Therapeutics Inc. Q3 2025 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: September 3, 2025 Pelthos Therapeutics Inc.

By: /s/ Francis Knuettel II

Name: Francis Knuettel II
Title: Chief Financial Officer





#### 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 for Zesluvmi, timing, progress and results of any preclinical and clinical trials, its estimates regarding the potential market opportunity for Zelsuvmi, its ability to develop its pipeline, its ability to protect its intellectual property and enforce its intellectual property rights, 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 ability to establish its market development capabilities to commercialize its products and generate any revenue, and the Company's ability to maintain regulatory approval of Zelsuvmi.

Forward-looking statements are provided to allow potential investors the opportunity to understan 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.



# **Corporate Profile**

Pelthos Therapeutics is a bio-pharmaceutical company committed to commercializing innovative, safe, and efficacious therapeutic products to help patients with unmet treatment burdens

- <u>Commercial launch</u> of first drug Zelsuvmi, for the treatment of Molluscum contagiosum ("MC") in July 2025
- First and only at home treatment for a large, underserved market treating contagious viral disease
  - ❖ 16.7 million affected people
  - ❖ Up to 6 million annual cases in the U.S.
  - Total addressable market worth in excess of \$20 billion at our WAC.
- Experienced management team with over 20 successful prior drug launches and continued growth, including Cosentyx, Otezla, Ohtuvayre, Xifaxan
- Upside option on NaV pain programs from predecessor
- Current peak Net Revenue forecast of \$175M per annum by 2028 and currently meeting or exceeding internal milestones

Key Data Points (as of 08/29/25, except where noted)			
Ticker	PTHS		
Stock Price	\$23.65		
O/S Shares of Common Stock (on an as converted basis)*	8.8M		
Market CAP	~\$205M		
Avg. Daily Trading Volume	59,000 shares		
Cash at close of merger	\$27.5M		
Investment to date	>\$400M		

#### **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



#### Frank Knuettel | Chief Financial Officer

- · 30 years of management experience in growing early-stage companies
- · Raised more than \$400 million via venture, public equity and debt offerings and managed more than 15 mergers and acquisition transactions along with large-scale licensing transactions with fortune 50 companies
- Holds numerous board positions, at both public and private companies, including Etheros Pharmaceuticals
   Earned an MBA from The Wharton School and a BA from Tufts University



#### 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



# **Board of Directors**















# Molluscum & Zelsuvmi Overview





#### **Molluscum Contagiosum**

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



#### Molluscum Contagiosum

is caused by a pox virus and is characterized by small, round, firm, umbilicated, often painless bumps. Even after healing, reinfection is possible if in contact with an infected person or object." ~17 million people infected in the U.S.



Point prevalence of 3-6 million children (ages 0-16) in the U.S.



MC can take up to five years to resolve without treatment<sup>2</sup>

There are **four known types** of MC virus
(MCVI, 2, 3, 4) with
MCVI being the most
common<sup>1</sup>



Peak incidence between 1-10 years of age. Up to **73% of children go untreated**<sup>3</sup>







# Zelsuvmi™ Has the Potential to Shift MC Treatment Paradigm

#### **Current Options**



 Other available topical treatment requires in-office visits every 3 weeks<sup>2</sup>



Painful, destructive treatments<sup>3</sup>



 Necessitates travel to HCP offices, adding to the time burden for MC patients and caregivers<sup>2</sup>



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



Breakthrough Product, Breakthrough Results

58.1%

Mean MC Lesion reduction count(1)

#### Zelsuvmi™

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

18, east-squares mean count reduction. See Figure 9. Browning 2C, Hebert A, Errice C, Castweight M, Meeda-Chubach T. Berdazimer Cell 1937s in a Clinically Meaningful Therapsube Intervence for Melascum Contagiosum. Abstract and poster presented at East Clinical 2014. Let Newsyn, NV. October 2-7-72, 7024. 2 [Litcherheld L F, Kwang P, C, Conzale MR, et al., Safety and Efficiency 4V-7-10. Exember 19, Conzale MR, et al., Safety and Efficiency 4V-7-70. Exember 19, Conzale MR, et al., Safety 19, S

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#### Zelsuvmi™ Efficacy Shown in Phase 3 Clinical Trials Drives Commercial Launch

#### Population

808 Males, 790 Females



Immunocompetent children and adults aged ≥6 months with 3-70 raised MC lesions

Mean age: 6.7 years (Range: 0.9 - 76.6 years)

# Intervention 1,598 participants randomized Zelsuvmi (terdazine) topical gel 10.3% 917 - Zelsuvmi<sup>TM</sup> Topical, once-daily application Topical, once-daily application

Topical, once-daily application of Zelsuvmi™ (berdazimer gel, 10.3%) to all active Mc lesions for up to 12 weeks

Topical, once-daily application of vehicle control gel to all active Mc lesions for up to 12 weeks

# Key Study Highlights Patients who applied Zelsuvmi<sup>M</sup>

Patients who applied Zelsuvmi™ for 12 weeks achieved a mean and median reduction in lesion count of 58% and 82%, respectively, compared to 36% and 43% for patients who applied a vehicle control gel



#### B-SIMPLE4 Study Locations



55 Clinics across the US

#### Safety

- Application site reactions were the most common adverse reaction associated with Zelsuvmi™
- Common application site reactions included mild pain and mild erythema (caused by increased blood flow)
- Minimal scarring incidences witnessed

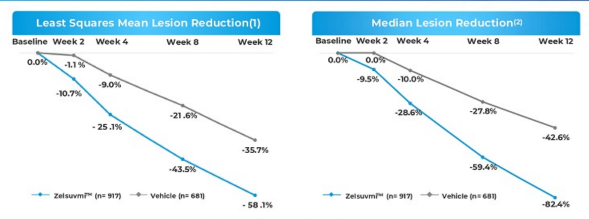
#### **B-SIMPLE4 Primary Outcome**

**32.4% of patients** treated with Zelsuvmi<sup>TM</sup> achieved complete clearance of MC lesions at week 12, compared to 19.7% of patients treated with vehicle control gel in the BSIMPLE-4 pivotal Phase 3 trial



### **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



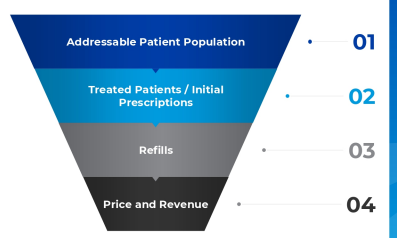
P<0.0001 at all time points, favoring Zelsuvmi™

§ Figure ® Browning 3C, Hebert A, Enlow C, Cartwright M, Meede-Chubechi T, Berdsülmer Cel 10.3% is a Clinically Meaningful Therapeutic Intervention for Meliuscum Contagiosum. Abstract and poster presented diffiel Clinical 2024. Las Vegas, Mr. October 24-27, 2004. 2] Figure 10. Browning IC, Hebert A, Enlow C, Cartwright M, Maeda-Chubachi T, Berdszimer Gel 10.3% is a Clinically Meaningful Therapeut diretereention for Meliuscum Contagiosum. Abstract and poster presented at Fall Clinical 2024. Las Vegas, NV. October 24-27, 2004.
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# Zelsuvmi Commercial Overview









#### **Key Leverage Points**

Up to 6 million new cases each year with an average untreated resolution time of 13 months, during which disease is highly contagious

Treated patients respond well to Zelsuvmi and experience dramatic reduction in lesion count and mean time to resolution

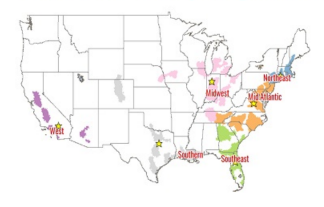
FDA approved for 12-week treatment regimen, but internal forecasts assume lower

WAC of \$1,950 per unit. Disclosed peak net revenues of \$175M per annum by 2028



#### **Commercial Launch Overview**

Sales Force of 50 Territory Managers Reaching Highest Molluscum Treaters



	NORTHEAST		MID ATLANTIC		SOUTHEAST
1.	Boston North, MA	1.	Toms River, NJ	1.	Atlanta North, GA
2.	Boston South, MA	2.	Philadelphia, PA	2.	Atlanta South, GA
3.	Providence, RI	3.	Baltimore, MD	3.	Jacksonville, FL
4.	Hartford, CT	4.	Washington, D.C.	4.	Fort Lauderdale, FL
5.	Long Island, NY	5.	Richmond, VA	5.	Orlando, FL
6.	Brooklyn, NY	6.	Raleigh, NC	6.	West Palm Beach, FL
7.	New York, NY	7.	Charlotte, NC	7.	Tampa, FL
8.	Summit, NJ	8.	Knoxville, TN	8.	Miami, FL
9.	Spring Valley, NY				
9.	MIDWEST		SOUTHERN		WEST
9.		1.	SOUTHERN New Orleans, LA	1.	WEST Phoenix North, AZ
1. 2.	MIDWEST	1.		1. 2.	
1.	MIDWEST Chicago North, IL		New Orleans, LA		Phoenix North, AZ
1. 2.	MIDWEST Chicago North, IL Chicago South, IL	2.	New Orleans, LA Houston North, TX	2.	Phoenix North, AZ Phoenix South, AZ
1. 2. 3.	MIDWEST Chicago North, IL Chicago South, IL Indianapolis, IN	2.	New Orleans, LA Houston North, TX Houston South, TX	2.	Phoenix North, AZ Phoenix South, AZ San Bernardino, CA Santa Ana, CA
1. 2. 3. 4. 5.	MIDWEST Chicago North, IL Chicago South, IL Indianapolis, IN Grand Rapida, MI	2. 3. 4.	New Orleans, LA Houston North, TX Houston South, TX Fort Worth, TX	2. 3. 4.	Phoenix North, AZ Phoenix South, AZ San Bernardino, CA Santa Ana, CA Los Angeles South, C
1. 2. 3. 4.	MIDWEST Chicago North, IL Chicago South, IL Indianapolis, IN Grand Rapids, MI Detrolt, MI	2. 3. 4. 5.	New Orleans, LA Houston North, TX Houston South, TX Fort Worth, TX Dallas, TX	2. 3. 4. 5.	Phoenix North, AZ Phoenix South, AZ San Bernardino, CA

Fully built out commercial team: Territory managers supported by Sales Training, Marketing, Commercial Operations & Market Access teams



# Zelsuvmi Tactical Launch Approach

#### **Driving Awareness and Adoption**

#### **Health Care Providers Education**













#### Consumer/Patient Education & Awareness













Advocacy & Partnerships

Zelsuvmi

ZELSUVMI GO Patient Support Program











Patient CRM Platform



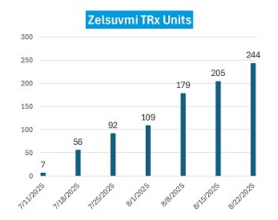




Patient Testimonial Videos



# Strong Uptake of Zelsuvmi in early Launch Phase



#### Zelsuvmi TRx & HCP Prescribers

	Week Ending	TRx Count	TRx Units	Prescribers
	7/11/2025	7	7	5
	7/18/2025	53	56	37
	7/25/2025	89	92	67
L	8/1/2025	109	109	71
	8/8/2025	176	179	126
	8/15/2025	199	205	150
	8/22/2025	234	244	162
	Total	867	892	*501

<sup>\*</sup> Total Unique Prescribers

Data Source: Symphony Health- Metys Data

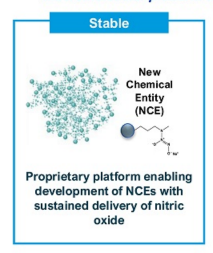
# Nitricil<sup>TM</sup> Platform & NaV1.7 Pipeline Overview

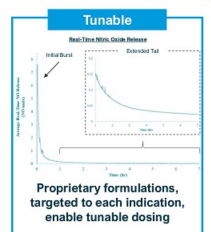


#### The Nitricil™ Platform



#### Macromolecular platform to achieve stable, tunable and druggable delivery of nitric oxide









# **Nitricil Platform Pipeline**

Asset Description	Asset Description	Approx Time to NDA Filing	Market Potential
SB204 (Acne)	Berdazimer topical gel, 3.4% for treatment of acne vulgaris. Phase 3 Clinical stage.	4.5 years	\$\$\$
SB414 (AD/Psoriasis)	Berdazimer topical cream, dose TBD, for treatment of mild to moderate atopic dermatitis. Phase 1/2 Clinical stage.	7.5 years	\$\$\$ (AD) \$\$ (Psoriasis)
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	\$



# NaVI.7 Pipeline

Product/ Indication	Asset Description	Approximate Time to NDA Filing	Market Potential
CT2000 Eye Drops Chronic Ocular Pain	CC8464 1%, 1.25% and 1.5% ophthalmic solution Phase 1-2a ready	3-4 years	\$8 billion globally
CT2000 Eye Drops Acute Ocular Pain	CC8464 1%, 1.25% and 1.5% ophthalmic solution Phase 1-2a ready	2-3 years	\$400 million globally
CT3000 depot Nerve Blocks	CC8464 5% and 10% depot injectable Preclinical Stage	5+ years	\$300-570 million globally
CC8464 Oral Erythromelalgia	CC8464 melt-granulation capsules 50mg, 100mg, 400mg Phase 1 stage	5+ years	\$2.4 billion globally
CC8464 Oral Small Fibre Neuropathy	CC8464 melt-granulation capsules 50mg, 100mg, 400mg Preclinical stage	5+ years	\$50 million – 100 million
CC8464 Oral Acute Pain	CC8464 melt-granulation capsules 50mg, 100mg, 400mg Preclinical stage	5+ years	\$20 billion globally



# **Key Highlights**



Commercially Launched

Zelsuvmi™ was FDA approved In January 2024, and commercially launched in July 2025 as the first and only at-home treatment aimed to revolutionize how MC is treated today for patients ≥ 1 year old



Significant Unmet Need and Sizeable Market Opportunity

Large market potential, with up to 6M new cases annually. Treatment of new cases alone has a total addressable market potential of over \$20 billion



Zelsuvmi™ Differentiated Characteristics Zelsuvmi<sup>™</sup> is a topical gel that uses proprietary nitric oxide release technology and is applied once daily at-home with very good safety profile; opportunity to replace and complement current approved treatment options that are painful and require in-person visits



Strong, Proven Clinical Efficacy In the combined results from the three Phase 3 clinical trials, patients who applied Zelsuvmi™ for 12 weeks achieved a mean and median reduction in lesion count of 58% and 82%, respectively, compared to 36% and 43% for patients who applied a vehicle control gel



**Barriers to Entry** 

Pelthos' bespoke manufacturing processes require a dedicated line and manufacturing of API under extremely high pressures with stringent safety protocols and procedures; robust set of FDA Orange Book listed patents



Biopharmaceutical Platform Poised for Growth Pelthos is strategically positioned to execute and integrate complex, synergistic acquisitions, serving as a platform for investors seeking a strong foothold in the specialty biopharmaceutical market



**Financial Opportunity** 

Retained Channel Therapeutics' pipeline of several NaVI.7 programs following business combination, providing further upside optionality. Currently exploring best monetization strategies.





#### Contacts

Investor Inquiries: Mike Moyer LifeSci Advisors, LLC Managing Director mmoyer@lifesciadvisors.com

Media: KWM Communications Kellie Walsh / Rachel Kessler pelthos@kwmcommunications.com (914) 315-6072



#### Pelthos Therapeutics to Participate in the 2025 Wells Fargo Healthcare Conference

Management's presentation will take place on Wednesday, September 3<sup>rd</sup>, 2025 at 1:30 pm (ET)

DURHAM, N.C., Sept. 2, 2025 (GLOBE NEWSWIRE) — Pelthos Therapeutics Inc. (NYSE American: PTHS), a biopharmaceutical company ("Pelthos" or the "Company") committed to commercializing innovative therapeutic products for high unmet patient needs, today announced that Scott Plesha, CEO, and Frank Knuettel, CFO, will participate in the Wells Fargo Healthcare Conference. Management's presentation will take place on Wednesday, September 3<sup>rd</sup>, 2025 at 1:30 p.m. E.T.

For one-on-one meeting requests please contact your institutional sales representative. A webcast of the presentation will be available on the Events and Presentations pages of Pelthos' website at <a href="https://pelthos.com">https://pelthos.com</a>. Archived replays will be available for 90 days following the event.

#### **About Pelthos Therapeutics**

Pelthos Therapeutics is a biopharmaceutical company committed to commercializing innovative, safe, and efficacious therapeutic products to help patients with unmet treatment burdens. The company's lead product ZELSUVMI<sup>TM</sup> (berdazimer) topical gel, 10.3%, for the treatment of *molluscum contagiosum*, was approved by the U.S. Food and Drug Administration in 2024. More information is available at <a href="https://www.pelthos.com">www.pelthos.com</a>. Follow Pelthos on <a href="https://www.pelthos.com">LinkedIn</a> and <a href="https://www.pelthos.com">X</a>.

#### Contacts

#### **Pelthos Investor Inquiries:**

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#### Media:

KWM Communications Kellie Walsh / Rachel Kessler pelthos@kwmcommunications.com (914) 315-6072