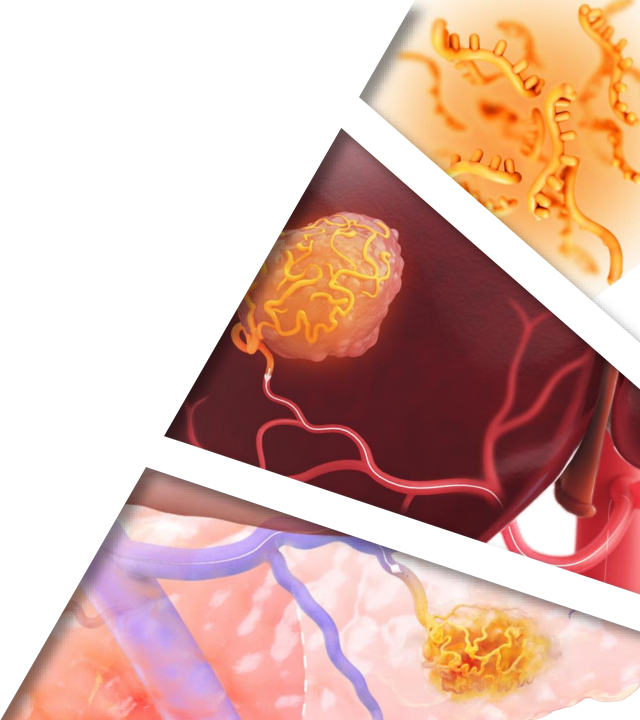


TriSalus Life Sciences

January 2025



Disclaimer

Certain statements in this presentation may constitute "forward-looking statements" within the meaning of applicable United States federal securities laws. Forward-looking statements include, but are not limited to, statements regarding TriSalus's expectations, hopes, beliefs, intentions or strategies regarding the future, including, without limitation, statements regarding TriSalus's business strategy and clinical development plans; the safety and efficacy of TriSalus's product candidates; TriSalus's plans and expected timing concerning clinical trials, clinical trial enrolment and clinical trial results; the size and growth potential of the markets for TriSalus's products and TriSalus's ability to serve those markets; TriSalus's ability to compete with other companies; TriSalus's expected financial results as of and for the year and quarter ended December 31, 2023; TriSalus's projected financial results and expected cash runway; TriSalus's ability to partner with other companies; and TriSalus's products continuing to be subject to a favorable reimbursement environment. In addition, any statements that refer to projections, forecasts, or other characterizations of future events or circumstances, including any underlying assumptions, are forward-looking statements. The words "anticipate," "continue," "could," "estimate," "expect," "may," "might," plan," "possible," "potential," "predict," "project," "should," "strive," "would" and similar expressions may identify forward-looking statements, but the absence of these words does not mean that statement is not forward-looking. Forward-looking statements are predictions, projections, and other statements about future events based on current expectations and assumptions and, as a result, are subject to risks and uncertainties.

Such statements are subject to a number of known and unknown risks, uncertainties and assumptions, and actual results may differ materially from those expressed or implied in the forward looking statements due to various important factors, including, but not limited to: changes in business, market, financial, political and legal conditions; unfavorable changes in the reimbursement environment for TriSalus's products; TriSalus's product candidates not achieving success in preclinical or clinical trials or not being able to obtain regulatory approval, either on a timely basis or at all; future clinical trial results/data may not be consistent with interim, initial or preliminary results/data or results/data from prior preclinical studies or clinical trials; TriSalus's ability to maintain and grow its market share; the size of the addressable markets for TriNav and TriSalus's product candidates being less than TriSalus estimates; TriSalus's ability to successfully commercialize any product candidates that are approved; TriSalus's ability to continue to fund preclinical and clinical trials for its product candidates; future economic and market conditions; the effects of competition on TriSalus's business; risks relating to the uncertainty of the projected financial information with respect to TriSalus; the ability of the company to raise money to finance its operations in the future; and the outcome of any potential litigation, government and regulatory proceedings, investigations and inquiries. You should carefully consider the risks and uncertainties described in the "Risk Factors" section of TriSalus's Quarterly Report on Form 10-Q for the quarter ended September 30, 2023, and other documents filed by TriSalus from time to time with the SEC. These filings identify and address other important risks and uncertainties that could cause actual events and results to differ materially from those expressed or implied in the forward-looking statements. Forward-looking statements, whether as a result of new information, f

Certain financial information and data in this presentation may be unaudited and may not conform to Regulation S-X promulgated under the Securities Act of 1933, as amended. Accordingly, such information and data may not be included, adjusted, or presented differently in any documents filed with the SEC.



Improving Therapeutic Delivery to Liver & Pancreatic Tumors

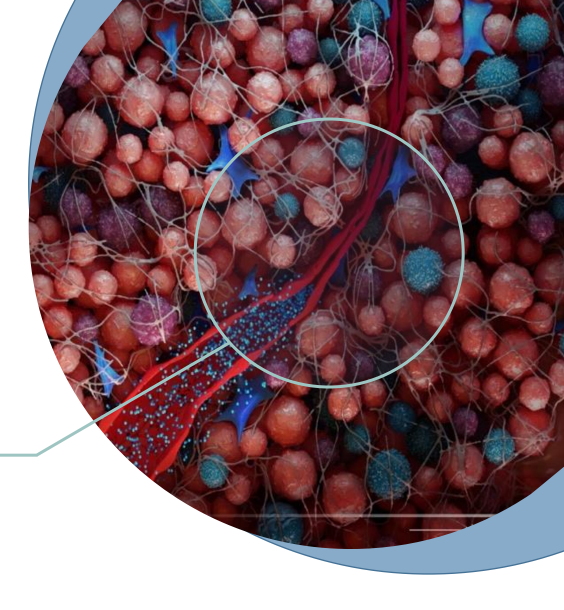
Core Med Tech Business	 A high-growth MedTech business with strong commercial potential, significant upside from expansion in existing and new applications
Our Technology	 Pressure Enabled Drug Delivery™ (PEDD™) infusion, which improves therapeutic delivery for hepatocellular carcinoma, pancreatic cancer, and other solid liver tumors Nelitolimod, a TLR9 agonist, delivered via PEDD to liver and pancreatic tumors
Attractive Markets	 PEDD market opportunity exceeds \$1 billion Nelitolimod (SD-101), if approved for combination with locally advanced pancreatic cancer, adds \$1 billion upside
Upcoming Milestones	 Nelitolimod, a TLR9 agonist, reverses immunosuppression and demonstrated Phase 1 proof of concept in Uveal Melanoma Liver Metastases. Phase 1 in Locally Advanced Pancreatic Cancer completing enrollment
Additional Upside	Additional growth from new product launches, pipeline advancement anticipated over next 18 months
2025 Outlook	 Anticipate 2024 revenue of ~\$28-30MM, with capacity to sustain 50% annual growth rate in 2025 © 202 TriSalus™ Life Sciences. All Rights Reserved. Strictly Confidential. Not for Distribution.

Tumor microenvironment limits drug delivery in liver and pancreatic tumors

- High intra-tumoral pressure in solid tumors limits efficient drug delivery to tumor
- Elevated interstitial fluid pressures reduce movement of fluid from vessel into tissue
- Lymphatic system within tumors is often underdeveloped and cannot drain fluids away

Limited drug uptake due to collapsed vessel

(<1% in some settings with IV delivery)



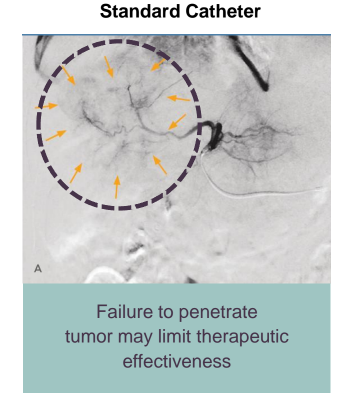
REFERENCES: 1. Kiet al. "Measurement of Tumor Pressure and Strategies of Imaging Tumor Pressure for Radioimmunotherapy." Nuclm, Hyeon-Gi ear medicine and molecular imaging vol. 53,4 (2019): 235-241. doi:10.1007/s13139-019-00598-7. 2. Heldin et al, "High Interstitial Fluid Pressure An Obstacle in Cancer Therapy," Nature Review, Vol 4, Oct 2004. 3. Sheth RA, et al. J Vasc Interv Radiol. 2013;24:1201-1207.4. Jain RK, Stylianopoulos T. Nat Rev Clin Oncol. 2010;7(11):653-664. DOI: 10.1038nrclinonc.2010.139., Wilhelm et al. (2016) Nature Reviews Materials 1.5:16014.



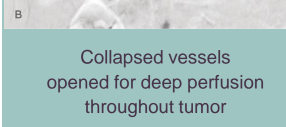
PEDD opens collapsed vessels, improving drug delivery in high pressure tumors¹

Angiogram of liver tumor vessels demonstrating PEDD method:

- Opens collapsed tumor vessels
 wider than conventional catheter
- Delivers increased contrast dye (surrogate for therapeutic) to tumor
- Protects normal tissue
 from chemo or radiation by reducing reflux



PEDD Method



Same patient several minutes apart

1. TriSalus images and data on file

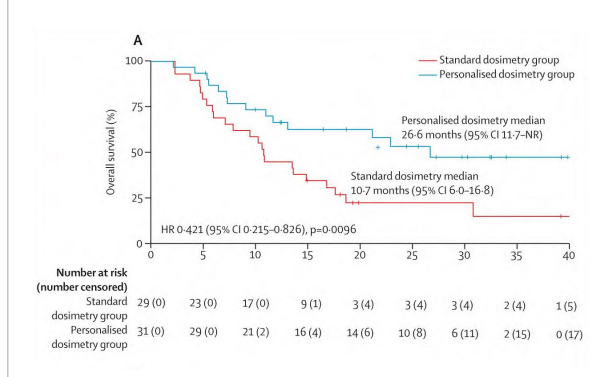


Increased delivery of Y90 beads to tumor correlated with improved survival

THE LANCET Gastroenterology & Hepatology

Personalised versus standard dosimetry approach of selective internal radiation therapy in patients with locally advanced hepatocellular carcinoma (DOSISPHERE-01): a randomised, multicentre, open-label phase 2 trial

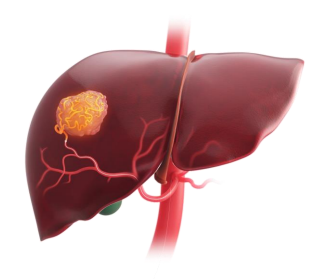
Etienne Garin*, Lambros Tselikas*, Boris Guiu, Julia Chalaye, Julien Edeline, Thierry de Baere, Eric Assenat, Vania Tacher, Corentin Robert, Marie Terroir-Cassou-Mounat, Denis Mariano-Goulart, Giuliana Amaddeo, Xavier Palard, Antoine Hollebecque, Marilyne Kafrouni, Hélène Regnault, Karim Boudjema, Serena Grimaldi, Marjolaine Fourcade, Hicham Kobeiter, Eric Vibert, Samuel Le Sourd, Lauranne Piron, Danièle Sommacale, Sophie Laffont, Boris Campillo-Gimenez, Yan Rolland, on behalf of the DOSISPHERE-01 Study Group†



Garin, E. et al. Personalised versus standard dosimetry approach of selective internal radiation therapy in patients with locally advanced hepatocellular carcinoma (DOSISPHERE-01): a randomised, multicentre, open-label phase 2 trial. Lancet Gastroenterol. Hepatol. 6, 17–29 (2021)



TriNav® Infusion System: a better solution for drug delivery



TriNav Infusion System

Commercial-stage, FDA-cleared technology using the proprietary PEDD method

510(k) cleared device for vascular access throughout the body except the heart and brain

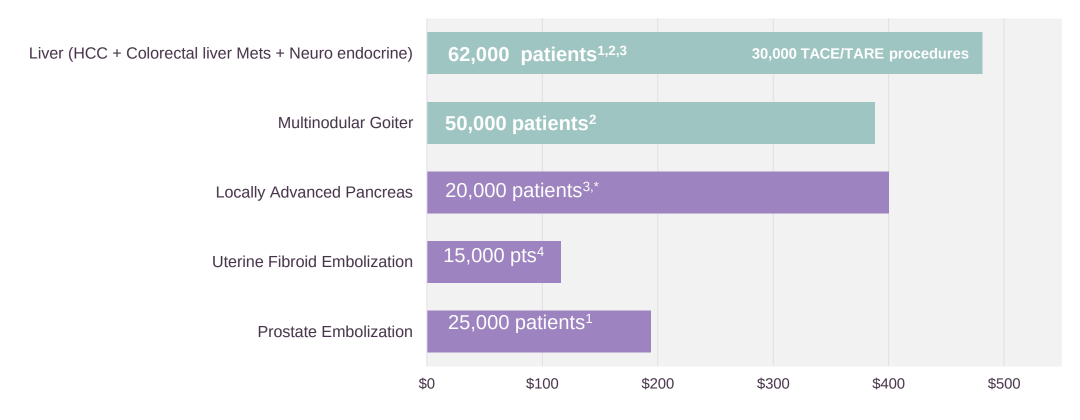
Unique HCPCS reimbursement code for procedures using the TriNav system

Validated in multiple clinical and HEOR studies

Addressable market in excess of \$1.5B



Combined U.S. PEDD total addressable market is >\$1.5 billion annually



■ Current Market - \$ 900MM
■ Additional Market - \$ 700MM

Source: 1. American Cancer Society, National Cancer Institute SEER Database, Horn, Epidemiology of Liver Metastases, Cancer Epidemiology, 2020, TriSalus Assumption; 2.https://my.clevelandclinic.org/health/treatments/7016-thyroidectomy, Ho TW et al. Utilization of thyroidectomy, Am J Surg 2011;201:570-4.; 3. American Cancer Society, National Cancer Institute SEER Database, TriSalus Assumptions 4. ACOG Committee Opinion #293, Obstetrics & Gynecology 103(2):p 403-404, February 2004.



Analysis of real-world data provides evidence that TriNav system successfully treats complex liver cancer patients¹

Population/Setting

- Retrospective analysis of 300 million patient claims over 3 years
- 98% of all payors
- Compared 258 TriNav patients to 8,940 conventional microcather patients

TriNav Patient Type: Key Findings

TriNav patients are more complex:

- More comorbidities and more liver-related adverse events
- More likely to have had prior embolization and/or prior systemic therapy
- Sicker and showed a higher burden of disease

Comparative Findings

- In chemoembolizations, TriNav delivered 40% more doxorubicin
- Higher disease burden patients receiving TriNav had outcomes similar to healthier non-TriNav patients
- In matched cohort analyses, TriNav patients did better
 - 48% increase in liver transplantation
 - 50% reduction in 30-day inpatient admissions
 - 17% reduction in complications
 - 40% reduction in fatigue

^{1.} Cook et al, Real-world evidence of Pressure-enabled Drug Delivery, Current Medical Research and Opinion, March 2024



Focused, accessible hospital market with attractive reimbursement

Highly Targeted Market

- ~400 hospitals cover 95% of procedure volume
- Targeting 450 out of the 1,000
 Interventional Radiologists
 who perform embolizations
- Commercial team partner to educate physicians and staff to drive adoption

Traditional Medicare Patients

- CMS issued HCPCS procedural code C9797, reimbursed under APC 5194 (Level 4 endovascular procedures), effective January 1, 2024
- C9797 exclusive to PEDD devices
- Reimbursement rate \$17,957 for 2025
- TriNav selling price \$7,750/catheter as of January 1, 2025

Private Payer Patients

- Commercial payers generally follow Medicare guidelines
- Payment rates generally 105% 125% of CMS payment rate



Technology pipeline: opportunities for further expansion

TriNav System
510k Cleared and
On Market

TriNav receives
Transitional
Passthrough Payment

Pancreas TriSalus Infusion System 510k Cleared TriNav 2.0

Launch Q1 2024

Currently in Phase 1 clinical trial with nelitolimod

Large Vessel
TriNav System
510k Cleared

Launched 2H 2024

Next Generation

Pancreatic Infusion

2.0 in Development

Integrated pressure measurement

Next Generation TriNav System with Integrated Sensing in Development

Integrated pressure measurement



Liver



Pancreas



SECTION 2

Future Areas of Growth

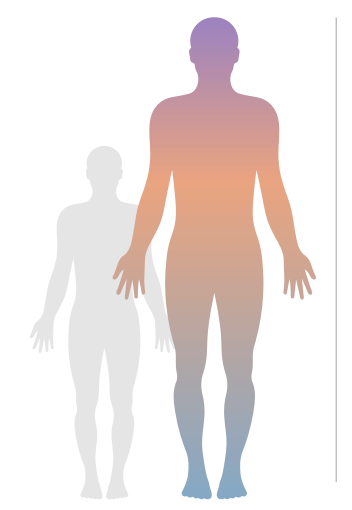
New applications and product launches



Data support for TriNav system in "complex patients"

Deliver Program initial focus is on four studies:

- PEDD benefits in multinodular goiter
- Registry study for uterine fibroid embolizations
- PEDD benefits in complex patients
- cTACE for colorectal and neuroendocrine liver metastases
- Potential expansion to additional populations
- Collaboration with leading IRs and Oncology KOLs



Multiple comorbidities

Previous embolization

Multi-focal disease – breast cancer liver mets

Hypovascular tumors colorectal cancer

Uterine fibroid

Large tumor size

Multinodular goiter



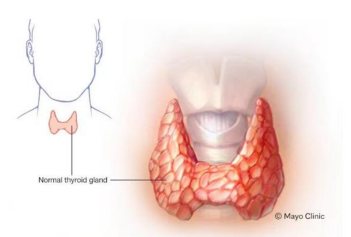
MultiNodular Goiter

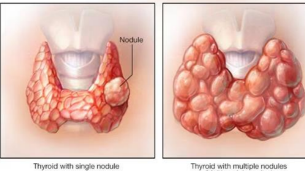
Novel minimally invasive approach via PEDD with potential to become standard-of-care



Multinodular goiter - large U.S. market opportunity

High unmet need in thyroid artery embolization with continued expected growth





~5% of population affected

Risk factors

Iodine deficiency
Female sex
Metabolic syndrome

Thyroidectomy

Risks of laryngeal nerve injury, long-term hormone replacement therapy, bleeding

Treatment

Watchful waiting,

Medical therapy

Surgery (Thyroidectomy)

Radioactive iodine (RAI) therapy

Radiofrequency ablation

50,000

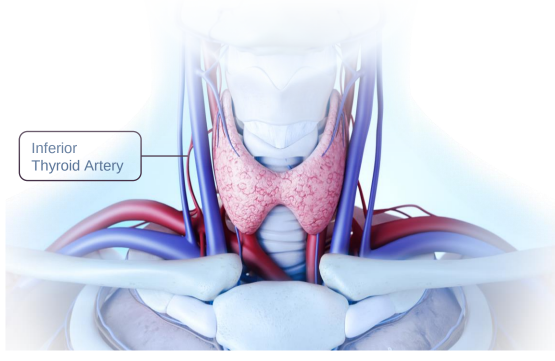
potential embolization patients per year

Opportunity: Offer alternative to surgery and avoid long-term thyroid replacement therapy



TriNav system has potential to be standard of care for treatment of multinodular thyroid disease

TriNav system enables treatment of the entire gland through the inferior thyroid arteries minimizing complication of stroke



Ablation	Surgery	Radioiodine Therapy	Embolization with PEDD
Limited to smaller lesions	Recurrent laryngeal nerve injury	Sialadenitis	Able to embolize all tumor sizes
Potential for skin and nerve injury	Long-term hormone replacement	Long-term hormone replacement therapy	Preserves thyroid gland
	Secondary malignancy	Secondary malignancy	No radiation or microwave exposure
	Large tumors/goiters require extensive procedure		Uses bland beads to restrict blood flow for high-level tumor necrosis



Novel pancreatic infusion technology for use with standard of care treatments (y90) and in combination with nelitolimod

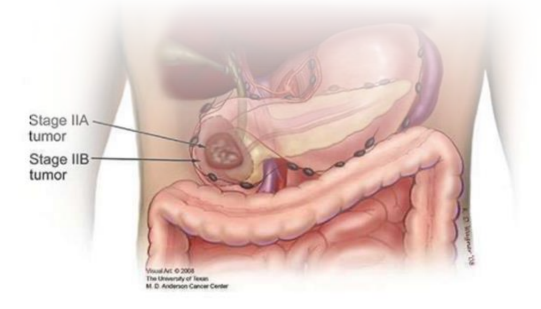
Locally Advanced Pancreatic Cancer



Large, growing market opportunity in locally advanced pancreatic cancer

High unmet need for improved treatments

Locally Advanced Adenocarcinoma



~30-50% of patient ineligible for surgery

Multi-agent chemotherapy primary treatment for most patients

Outcomes: 2L + overall survival approximately 5-6 months

25,000 potential patients per year

Opportunity: Deliver therapeutics to site of disease in combination with systemic therapy with incremental minimal toxicity

BMC Cancer volume 20, Article number: 203 (2020).



TriSalus developed a separate, novel PEDD method for the pancreas

FDA-cleared And In Phase 1 Clinical Trials With Nelitolimod

Poor blood flow limits drug access to pancreas^{1,2,3}

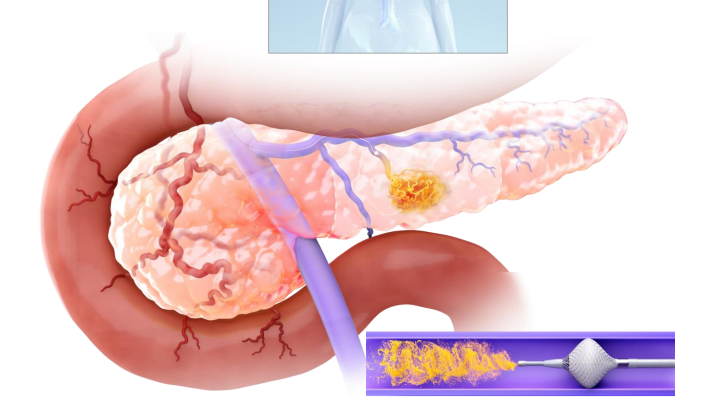
Pancreatic arteries difficult to access^{4,5}

Innovative retrograde venous approach eliminates need for balloons that eliminate blood flow^{6,7}

Target vessel pressure monitoring for safety, efficacy, and consistency

510k cleared

Phase 1 locally advanced pancreas data from MDACC was presented at SITC 2023



Rakesh Jain (2013) Normalizing Tumor Microenvironment to Treat Cancer: Bench to Bedside in Biomarkers. 31:17 2205-2218.
 DUED et al. Identified Pressure in Pagneratio Purcha Adaptography and Indigendated by a Gel-Eluid Phase. Biombusical Journal 110, 2106-2119.



19

DuFort et al, Interstitial Pressure in Pancreatic Ductal Adenocarcinoma Is Dominated by a Gel-Fluid Phase. Biophysical Journal 110 2106-2119.
 Soltani et al Numerical Modeling of Fluid Flow in Solid Tumors. PLoS ONE 6:6 e20344

Homma, H. et al. Cancer 89, 303–313 (2000).
 Rosemurgy, A. S. et al. J Pancreat Cancer 3, 58–65 (2017).

^{6.} Piras, C., Paulo, D. N. S., Paulo, I. C. A. L., Rodrigues, H. & Silva, A. L. da. Acta Cirurgica Brasileira 25, 105–110 (2010).
7. Moody, A. R. & Poon, P. Y. American Journal of Roentgenology 158, 779–783 (1992). 5. Okahara, M. et al. Abdom Imaging 35, 134–142 (2010)

Collaboration with Y90 partner for pancreatic infusion of Y90 beads

Pre-clinical

In normal swine (n=12, 6 head and 6 body)

25, 100, and 500 Gy to head or body of pancreas (2 per dose level for head and body/tail)

PET/CT for pancreatic dosimetry and SPECT/CT for off-target dose within 24 hours

Further dose ranging if deemed necessary

Phase 1- Y90 Delivered Pancreatic Infusion Device

To be Determined

Phase 2 - Concept & Regulatory Strategy

To be Determined

Summary of Objectives Met and Goals for Further Development

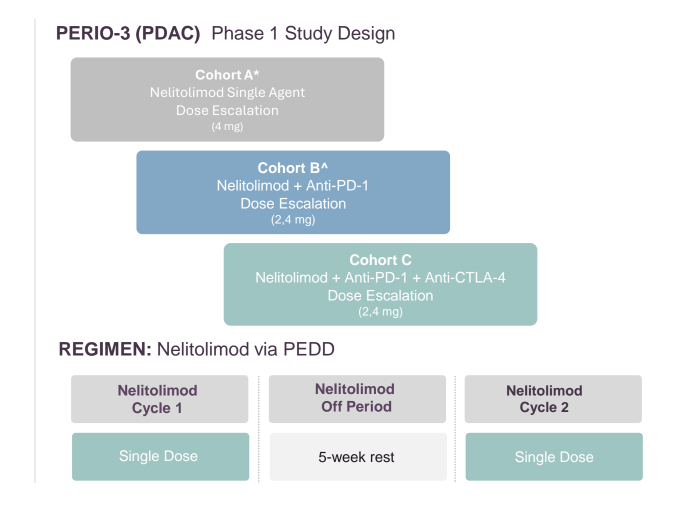
- Locally Advanced PDAC 1L+
- Cohort 1 3 dose levels, 3 per level
- Cohort 2 pancreatic head, 3 dose levels, n=3 per level
- Primary endpoint: safety and dose determination
- Secondary endpoint: progression free survival (PFS) and duration of disease
- PET/CT to quantify dose delivered target pancreatic tissue, non-target pancreatic tissue, and non-target extrapancreatic tissue
- Y90 localization to target and non-target tissue on PET/CT
- Extrapancreatic radiographic changes
- Necroscopy to examine target tissue and non-target tissue for radiation effects
- The combination of pancreatic infusion device and radiation has potential to improve response rates and reduce toxicity



20

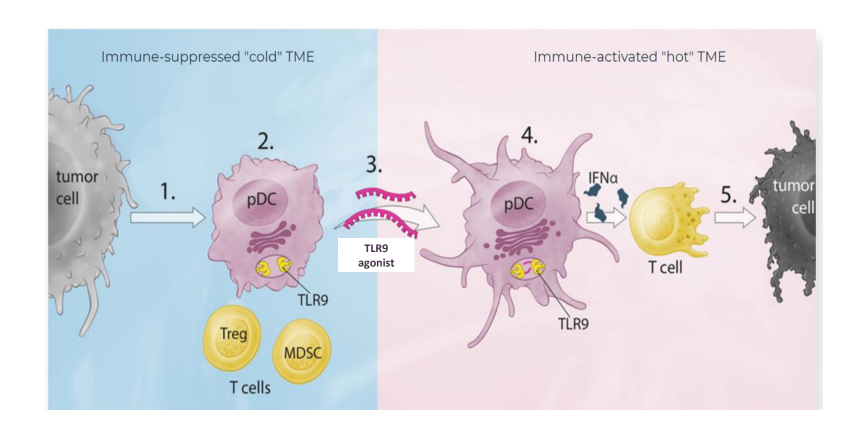
PERIO-3: nelitolimod + PEDD method for locally advanced pancreatic carcinoma

- Phase 1 trial currently enrolling at MD Anderson Cancer Center
- First time use of novel device with PEDD method to infuse into pancreatic tumors via venous vasculature
- Completed minimum enrollment at initial three dose levels without any safety dose limiting events
- Comparison of pre- and post-nelitolimod infusion PDAC tumor specimens revealed decreases in MDSC-associated genes and increased T cell activation
- Nanostring and flow cytometry analysis indicating systemic immune activation





Nelitolimod dual mechanism of action overcomes immunosuppression



- Nelitolimod binds to TLR9 receptors in the tumor
- 2. PEDD method allows for high concentration of therapeutic into the tumor
- 3. Nelitolimod is taken into tumor cells and eliminates immunosuppressive cells
- Nelitolimod activates immune system in the tumor



22

Clinical proof-of-concept

Dual moa with potential to enhance checkpoint activity both in tme and systemically

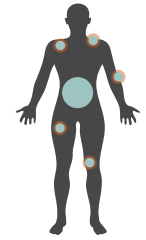
Tumor Pre-Nelitolimod

Responsive "Hot" **Tumor Post-Nelitolimod**

Enhanced immune Effects in Target Organ and Systemically







Systemic immune stimulation as evidenced by declines in ctDNA with disease control and early evidence for PFS²

MDSC accumulation

MDSC depletion

T cell infiltration

Enhanced chemotherapy, CPI/chemo combination activity

PEDD method shown to unlock dual MOA in liver and



T cell paucity

Immunotherapy failure

MDSC – myeloid derived suppressor cells. 1.Data on File. 2.Patel SITC 2023 pancreas¹

TS-PERIO-03 Phase 1 study for locally advanced PDAC

Phase 1 Single-agent Safety

Enrollment Completed







Dose escalation

Phase 1b Combination with Chemotherapy





Dose escalations with expansions to define Optimal Dose

Phase 2 - Concept & Regulatory Strategy

To be determined

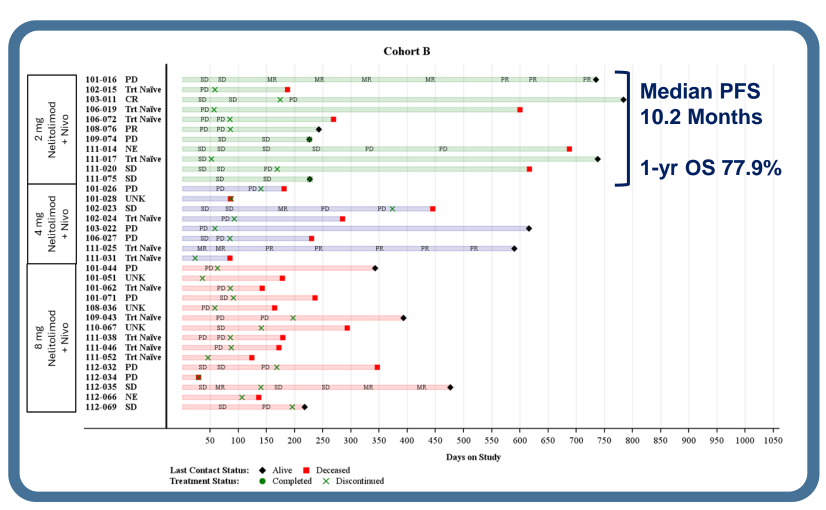
▲TriSalus[™]

Summary of Objectives Met and Goals for Further Development

- Enrolled 13 patients (0.5, 2, and 4 mg), mostly 3L and 2 patients 2L
- Awaiting data to mature
- Potential for breakthrough designation

24

Nelitolimod: SITC 2024 Phase 1 durable disease control and PFS in uveal melanoma liver metastases (UMLM)





- 67 patients and 69% were pre-treated (45% ICI, 18% tebentafusp)
- Grade 3/4 treatment-related AEs (TRAEs) in 13% of pts
- Recommended phase 2 dose was 2 mg nelitolimod + ICI (nivo or ipi/nivo, n=23)
 - 1-year OS = 74.7% (median, 20.6 mos)
 - 1-year PFS = 47.6% (median, 8.7 mos)
 - Disease control rate = 65%
 - OS similar in ICI-refractory (n=6, 80%) and ICI-naïve (n=17, 71%)
 - OS and PFS outcomes not dependent on HLA-A02:01 status
- o ctDNA clearance rate was 50%, with 76.9% responding (n=26)
 - 91% showing decrease at the 2 mg dose level (n=11)
 - ctDNA clearance correlated with OS in mUVM²
- Clinical benefit was associated with reduced tumor MDSCs and increased circulating IL-15/IL-18

Carvajal. SITC 2024. Rodrigues. Nature Communications. 2024

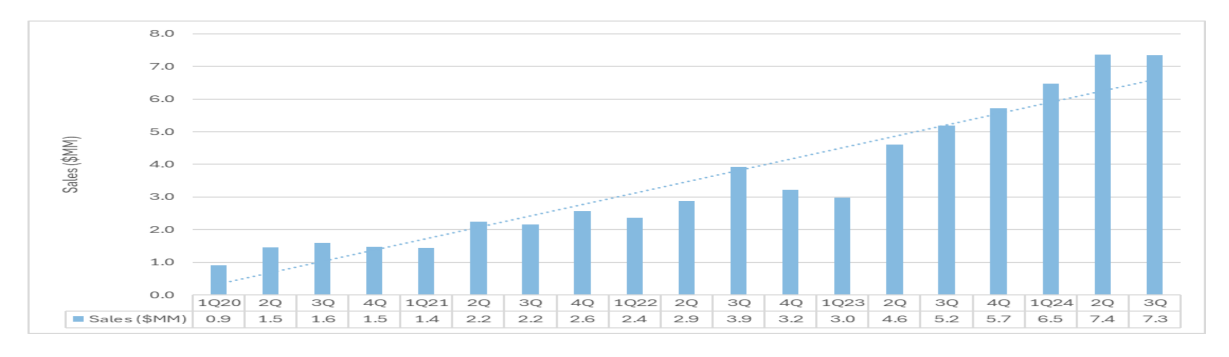


Corporate Update

Q3 2024



Strong multi-year growth forecasted



50% CAGR Projected Beyond 2024





Q3 2024 Highlights

Capitalization	TLSI (NASDAQ)	
Shares Outstanding ¹	30.5M	
Warrants Outstanding ²	7.3M	
52 Week Low – High ³	\$3.32 - \$10.42	
30d Average Daily Volume ⁴	41K	

42% total revenue growth over Q3 2023

Gross margins YTD of 86%

Cash and Investments of \$11.3M at September 30, 2024

Total debt of \$25M as of Q2 2024 with \$25M additional draws assumed in 2025

Cash runway through the end of 2025



2024 and 2025 key milestones

Catalyst	Indication	Timing
PROTECT	Launch of Pressure-enabled Retrograde Occlusive Therapy with Embolization for Control of Thyroid disease clinical study	Launched
Launch of TriNav Large	Hepatocellular Cancer, Uterine Fibroid and Liver metastases	Launched
HEOR TriNav DATA	2 nd year data comparing TriNav use in Complex patients in 300 mm patient claims	1H 2025
Launch of TriNav 2.0	Enhanced version of TriNav	1H 2025
Initiation of Phase 1 of Pancreatic Infusion with Y90	Phase 1 Locally Advanced Pancreatic patients with Y90	1H 2025
Launch of TriNav Thyroid Embolization	Full launch of TriNav Thyroid Embolization and Registry Data release	2H 2025
PERIO-3 Phase I Data	Release of PERIO-1 Phase 1 data	2H 2025



Veteran Industry Leadership















Mary Szela
CEO & President

Jim Young Chief Financial Officer

Sean Murphy
Chief Manufacturing, Strategy
& Business Development
Officer

Richard Marshak, VMD Chief Commercial Officer

Jennifer StevensChief Regulatory Officer

Jodi DevlinChief of Clinical Strategy &
Operations

Bryan Cox, PHD
Chief of Research









Baxter



























Improving Therapeutic Delivery to Liver & Pancreatic Tumors

We are	A high-growth MedTech business with strong commercial potential, significant upside from expansion in existing and new applications
Our Technology	 Pressure Enabled Drug Delivery™ (PEDD™) infusion, which improves therapeutic delivery for hepatocellular carcinoma, pancreatic cancer, and other solid liver tumors
Attractive Markets	 PEDD market opportunity exceeds \$1 billion Nelitolimod (SD-101), if approved for combination with locally advanced pancreatic cancer, adds \$1 billion upside
Upcoming Milestones	 Nelitolimod, a TLR9 agonist, reverses immunosuppression and demonstrated Phase 1 proof of concept in Uveal Melanoma Liver Metastases. Phase 1 in Locally Advanced Pancreatic Cancer completing enrollment
Additional Upside	Additional growth from new product launches, pipeline advancement anticipated over next 18 months
2025 Outlook	 Anticipate 2024 revenue of ~\$28-30MM, with capacity to sustain 50% annual growth rate in 2025 © 202 TriSalus™ Life Sciences. All Rights Reserved. Strictly Confidential. Not for Distribution.



Thank you

Contacts

James Young, SVP Investor Relations/Treasurer James.young@Trisaluslifesci.com

LifeSci Advisors

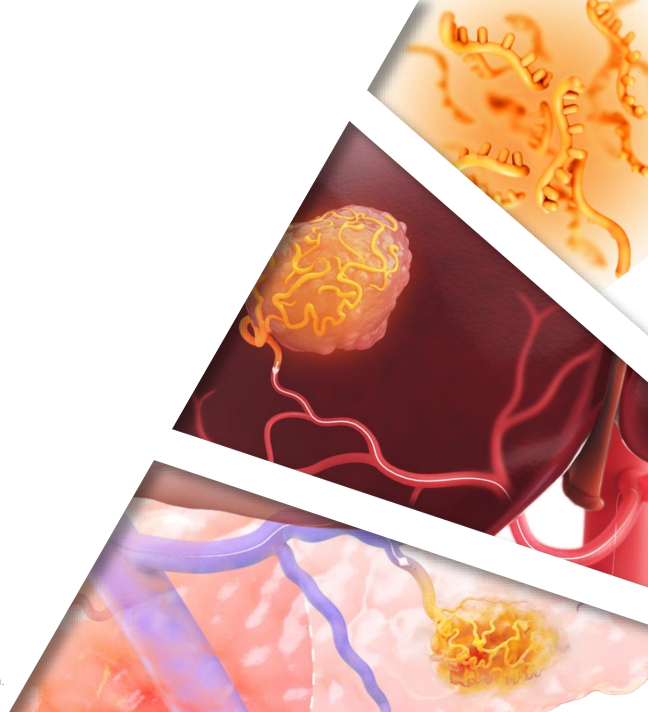
Jeremy Feffer, Managing Director jfeffer@lifesciadvisors.com





Thank You

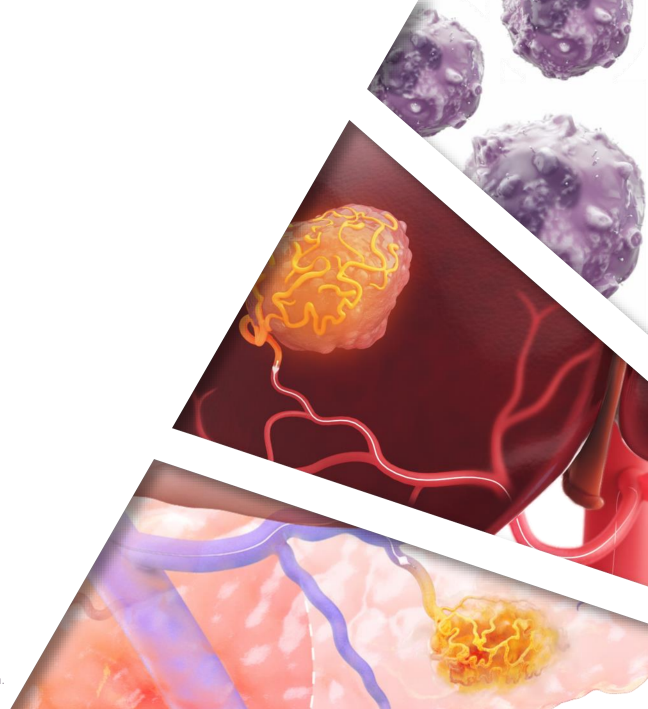
trisaluslifesci.com





Thank You

trisaluslifesci.com



Color Palette

In General, Use Only the Colors on the Top Row; Copy and Paste One of These Boxes to Start

Use the top row of these 6 colors for graphics, use Accent 1 & 2 first Top Color Top Color is is for Text for Subheads **Duplicate** White/Greys Accent 1 Accent 3 Accent 4 Accent 5 Accent 6 Accent 2 White SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE Text SAMPLE **SAMPLE SAMPLE** SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE **SAMPLE** SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE **SAMPLE** SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE SAMPLE **SAMPLE** SAMPLE SAMPLE

