

RENOVO | RX



Investor Presentation

Delivering therapy where it matters®

September 2026

NASDAQ | RNXT

Cautionary Note Regarding Forward-Looking Statements

This presentation and statements of the Company's management made in connection therewith contain forward-looking statements within the meaning of Section 27A of the Securities Act of 1933, and Section 21E of the Securities Exchange Act of 1934, including but not limited to statements regarding (i) the potential of RenovoCath® or TAMP™ as standalone commercial products, the estimated maximum total annual sales and addressable market for RenovoCath® and our commercialization plans in general, (ii) the prospects of our marketing and sales strategies, (iii) our clinical trials and studies, including the overall timing and timing for additional interim data readouts and patient enrollment for our ongoing TIGeR-PaC Phase III clinical trial study in LAPC, (iv) the potential for our product candidates to treat or provide clinically meaningful outcomes for certain medical conditions or diseases and (v) our commercialization strategies utilizing our TAMP technology. Statements that are not purely historical are forward-looking statements. These statements may be identified using words such as “may,” “expects,” “plans,” “aims,” “anticipates,” “believes,” “forecasts,” “estimates,” “intends,” and “potential,” or the negative of these terms or other comparable terminology. Forward-looking statements are based upon our current expectations and beliefs regarding future events, many of which, by their nature, are inherently uncertain, outside of our control and involve assumptions that may never materialize or may prove to be incorrect. These may include estimates, projections and statements which are based on current expectations and assumptions that are subject to known and unknown risks and uncertainties that may cause actual results to differ materially and adversely from those expressed or implied by these forward-looking statements. These risks include, among others: (i) the risk that our exploration of commercial opportunities for our TAMP technology may not lead to viable, revenue generating or profitable operations; (ii) circumstances which would adversely impact our ability to efficiently utilize our cash resources on hand or raise additional funding; (iii) the timing of the initiation, progress and potential results (including the results of interim analyses and the timing for patient enrollment) for TIGeR-PaC and any other preclinical studies, clinical trials and our research programs; (iv) the possibility that interim results may not be predictive of the outcome of our clinical trials, which may not demonstrate sufficient safety and efficacy to support regulatory approval of our product candidate; (v) that the applicable regulatory authorities may disagree with our interpretation of the data and cause delays in research and clinical development plans and timelines, and the regulatory process for our product candidates; (vi) future potential regulatory milestones for our product candidates, including those related to current and planned clinical studies; (vii) our ability to use and expand our therapy platform to build a pipeline of product candidates; (viii) our ability to advance product candidates into, and successfully complete, clinical trials; (ix) the timing or likelihood of regulatory filings and approvals; (x) our estimates of the number of patients who suffer from the diseases we are targeting and the number of patients that may enroll in our clinical trials; (xi) the commercialization potential of our other product candidates, if approved; (xii) our ability and the potential to successfully manufacture and supply our product candidates for clinical trials and for commercial use, if approved; (xiii) future strategic arrangements and/or collaborations and the potential benefits of such arrangements; (xiv) our estimates regarding expenses, future revenue, capital requirements and needs for additional financing and our ability to obtain additional capital; (xv) the sufficiency of our existing cash and cash equivalents to fund our future operating expenses and capital expenditure requirements; (xvi) our ability to retain the continued service of our key personnel and to identify, and hire and retain additional qualified personnel; (xvii) the implementation of our strategic plans for our business and product candidates; (xviii) the scope of protection we are able to establish and maintain for intellectual property rights, including our therapy platform, product candidates and research programs; (xix) the pricing, coverage and reimbursement of our product candidates, if approved; and (xx) developments relating to our competitors and our industry, including competing product candidates and therapies. Information regarding the foregoing and additional risks may be found in the section entitled “Risk Factors” in documents that we file from time to time with the Securities and Exchange Commission. Forward-looking statements included herein are made as of the date hereof, and RenovoRx does not undertake any obligation to update publicly such forward-looking statements to reflect subsequent events or circumstances, except as required by law.

RenovoRx at a Glance

RenovoRx is a commercial-stage targeted oncology platform with two value drivers – 1) FDA-cleared, high margin medical device business scaling rapidly and 2) Fully enrolled pancreatic cancer PH3 trial with survival read-out in '27



Driving adoption of **FDA-cleared RenovoCath®** across **21 U.S. commercial cancer centers (as of 8/12/26)** and **growing**, accelerating market penetration and strengthening commercial momentum.



Continues to estimate that **the initial TAM for RenovoCath as a standalone device** could translate into an **~\$400 million peak annual U.S. sales opportunity** for RenovoRx. Over time, as the Company expands into additional solid tumor indications, the Company believes it could unlock more than \$1 billion in peak annual sales potential.



Developing combination therapies based on **proprietary Trans-Arterial Micro-Perfusion (TAMP™) platform**; awarded 3 FDA Orphan Drug Designations.



Phase III TIGeR-PaC achieved full enrollment in August 2026; final data anticipated in the second half of 2027. TIGeR-PaC is evaluating intra-arterial gemcitabine delivered via RenovoCath (known as “**IAG**”) in locally advanced pancreatic cancer (LAPC).

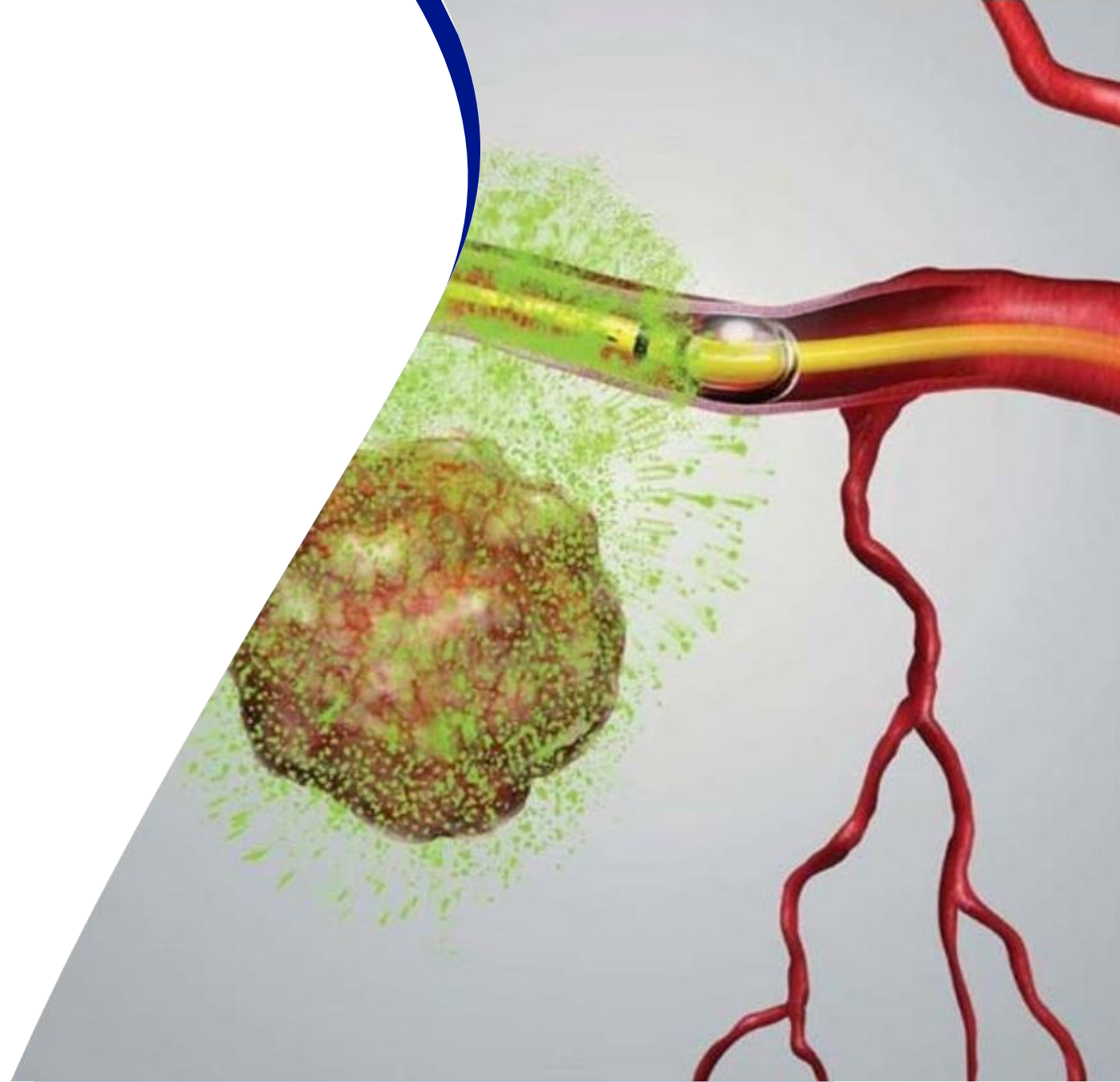


Led by **experienced Leadership** Team and Board of Directors.

¹ Based on Internal RenovoRx Estimates

Proprietary Trans-Arterial Micro-Perfusion (TAMP) Therapy Platform

Enabled by RenovoCath



Addressing a Significant Problem in Cancer Treatment



Hypervascular tumors are inadequately treated with current therapies

For example, liver tumors are highly vascularized

- Large tumor feeders = possible access for systemic therapy
- Feeders can be accessed and treated with current local therapy techniques
- Despite the abundance of blood vessels, systemic therapy may not always be effective due to drug delivery challenges
- Techniques like trans-arterial chemoembolization (TACE) and radioembolization are commonly used



Hypovascular tumors = major barrier to chemotherapy treatment success

Many tumors, like pancreatic tumors, bile duct tumors, etc. have poor blood supply

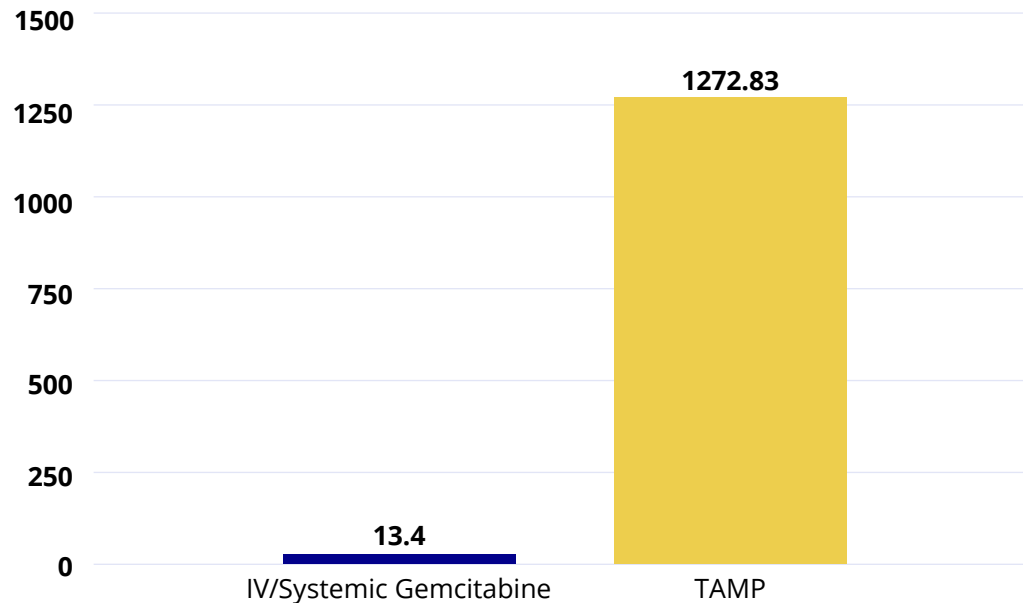
- No visible tumor feeder vessels
- Systemic therapy does not reach tumor tissue in high concentrations
- Inability to identify or engage tumor feeder vessels: local therapy ineffective
- Poor perfusion impacts drug delivery, leading to lower treatment efficacy

RenovoRx addresses this unmet medical need

TAMP Improves a Drug's Therapeutic Index

Increases Drug Concentration to Target Pathological Site by ~100X¹

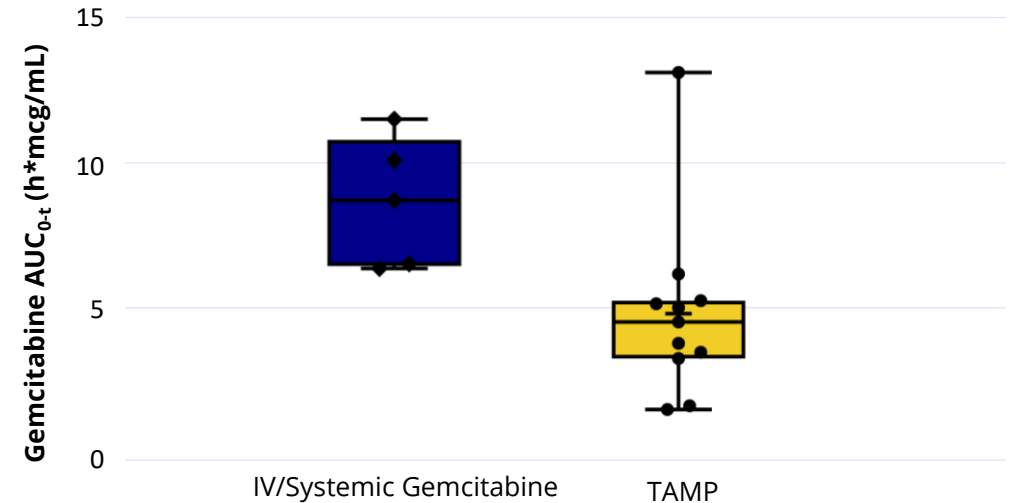
Tissue Concentration Gem (mcg/g)



¹Compared to IV Administration, as demonstrated in animal studies published in JVIR 2024 Article

Reduces Drug AUC_{0-t} by 43.5% Compared to IV Administration²

Drug Exposure IA vs. IV route



Impact of route of administration on drug exposure (area under the plasma concentration curve from time 0 to the last observable concentration [AUC_{0-t}]; h*mcg/mL; p=0.018)

²As demonstrated in Phase III TIGeR-PaC sub-study presented at ASCO-GI 2026

New PK data, presented at ASCO-GI 2026, from sub-study in Phase III trial strengthens scientific basis for TAMP as a targeted drug-delivery approach. These data show significantly lower systemic drug exposure compared to IV and provide further evidence that TAMP and IAG may offer the potential to deliver treatment more efficiently, improving local effectiveness while minimizing chemotherapy-related side effects.

TAMP

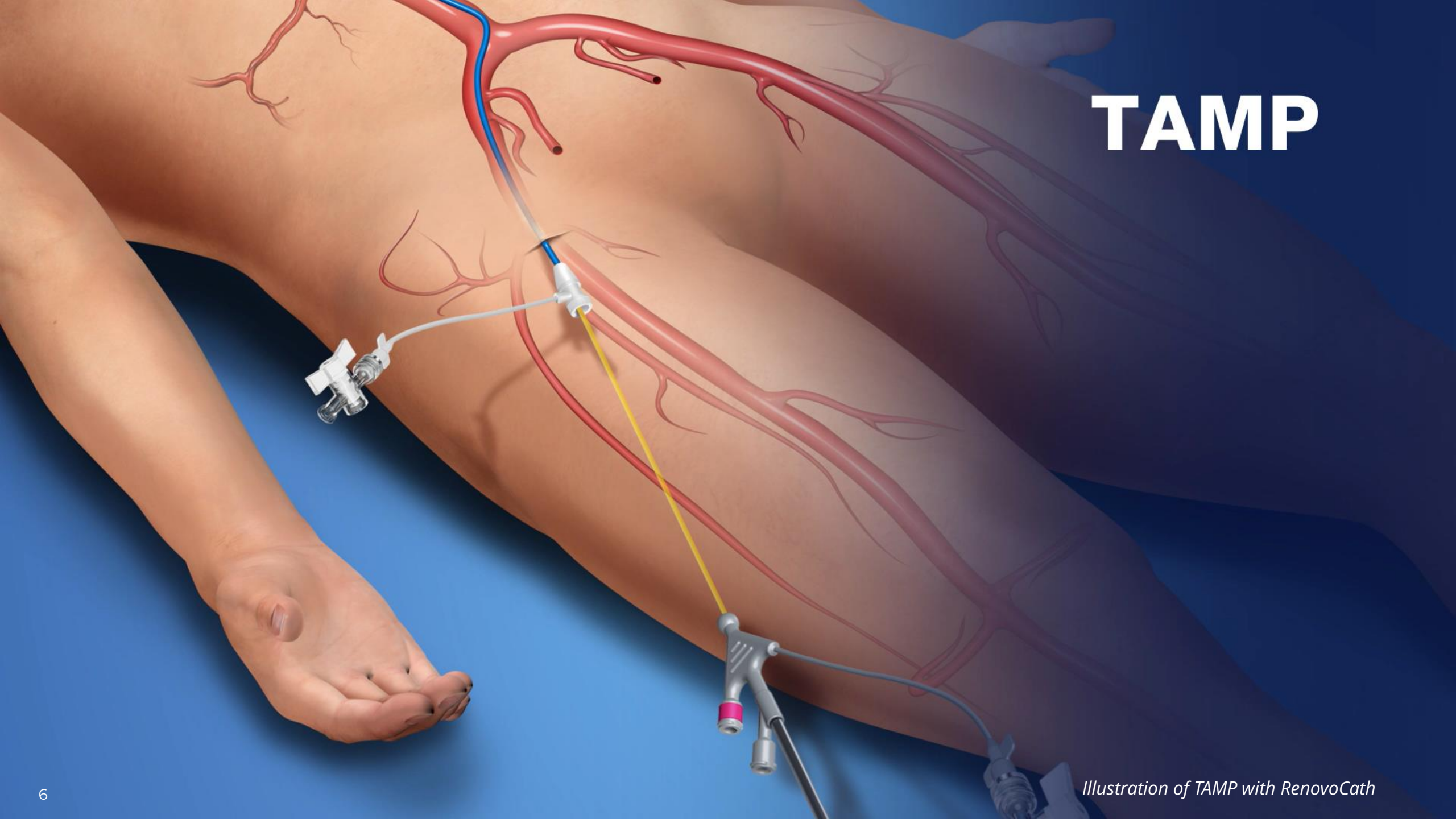
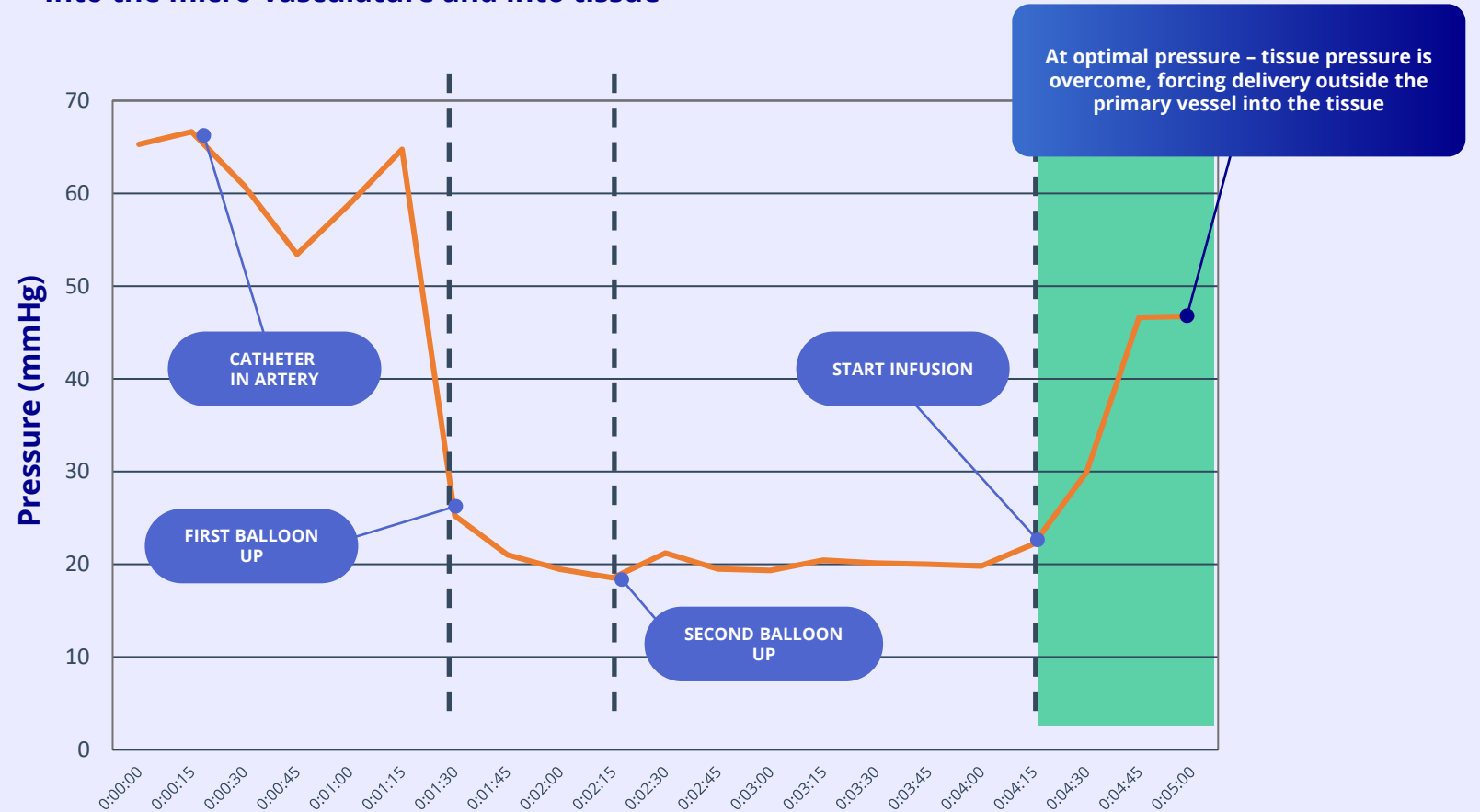


Illustration of TAMP with RenovoCath

Mechanism: Trans-Arterial Micro-Perfusion (TAMP)



Mechanism: after vessel isolation, increase in pressure forces drug across the artery wall into the micro-vasculature and into tissue



Source – RenovoRx Internal Animal Data presented at medical conferences.

Patient and Physician Experience

RenovoCath

RenovoCath Patient Experience

- ~8 treatments over 4 months (2x/month hospital visits) with disposable device (no implant)
- 20-minute infusion; ~90-minute outpatient procedure (shorter for subsequent procedures)
- Patients generally not put under general anesthesia (primarily conscious sedation for comfort)

RenovoCath Physician Experience

- Procedure easy to learn for interventional radiologists
- Transferable techniques utilized in liver-directed therapies resulting in fast learning curve for physicians
- Physicians demonstrate expertise after 2-3 proctored procedures and can train their colleagues

VS

Other Treatment Options

Other Patient Experience

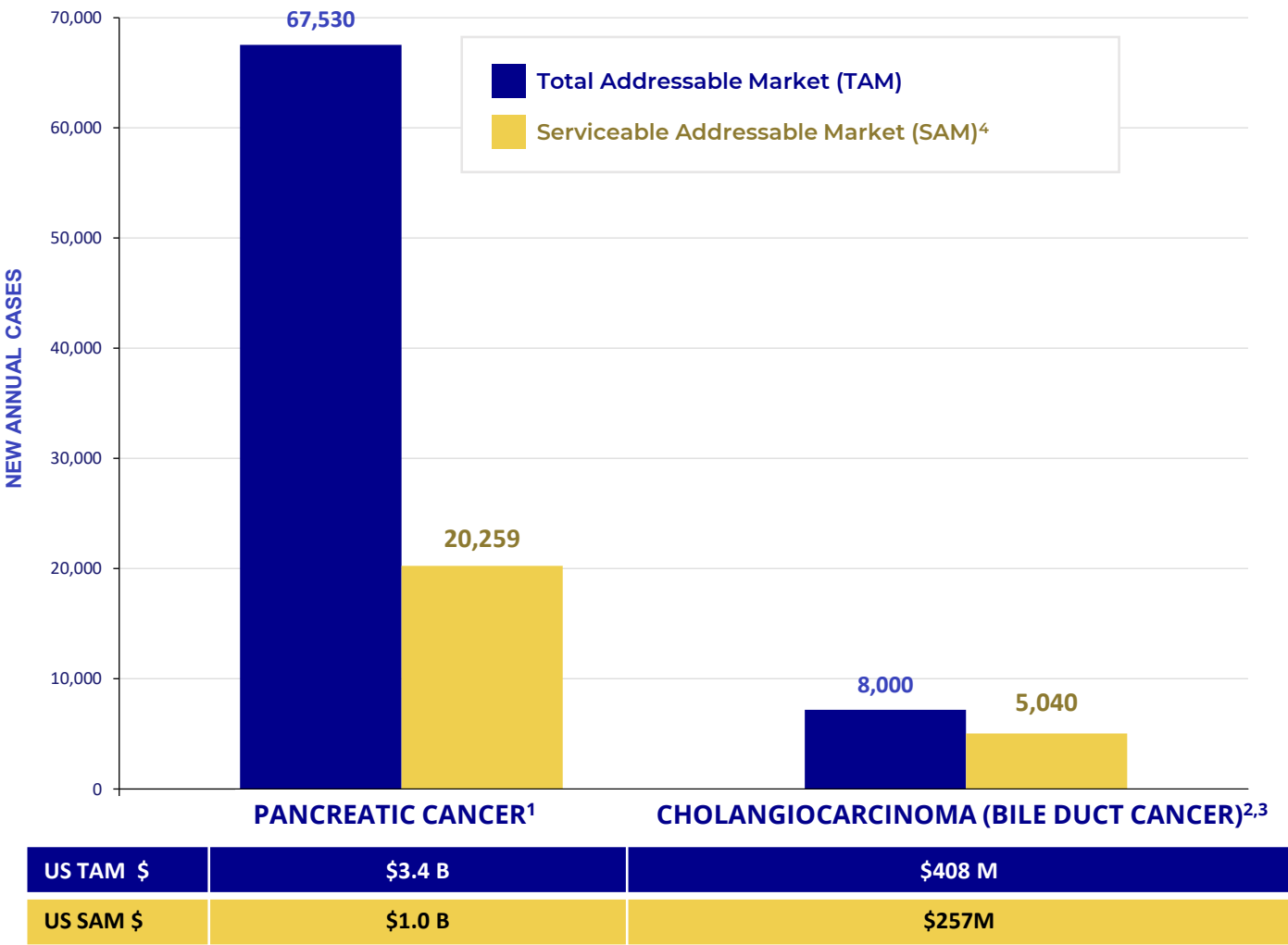
- Traditional systemic gemcitabine + Abraxane: 12 hospital/clinic visits over 4-month period or longer
- Other technologies may require overnight stay
- Other technologies may require general anesthesia
- Systemic chemo associated with days of lasting side effects

Other Physician Experience

- Majority of novel technologies require large sales/physician proctoring efforts with training courses and/or on-site support for every procedure

RenovoCath: A Drug-Delivery Platform Able to Serve Multiple Markets

INITIAL TARGETED ONCOLOGY APPLICATIONS IN UNITED STATES



FUTURE \$\$ GROWTH POTENTIAL

Future Expansion Opportunities Could Increase SAM:

- Borderline resectable pancreatic cancer
- Metastatic pancreatic cancer
- Sarcoma
- Non-small cell lung cancer
- Glioblastoma
- Pelvic tumors
- Others

1. seer.cancer.gov/statfacts/html/pancreas.html

2. <https://www.cancer.gov/pediatric-adult-rare-tumor/rare-tumors/rare-digestive-system-tumors/cholangiocarcinoma>

3. <https://pmc.ncbi.nlm.nih.gov/articles/PMC4492336/>

4. <https://curesarcoma.org/sarcoma-education/2026-sarcoma-statistics/>

5. Serviceable Addressable Market is Based on Percentages of RenovoRx's Internal Analysis of Total Addressable Market



Commercial Opportunity RenovoCath Standalone, FDA-Cleared Drug-Delivery Device

High Margin Large Market Opportunity for RenovoCath Standalone Device

RenovoCath US Market Opportunity

Analogous pressure-mediated delivery catheter ASPs	\$6k-\$8.5k/unit ¹
Annual average procedures	6 per patient ²
Peak annual revenue opportunity	\$400M ³

¹ <https://trisalulifesci.com/wp-content/uploads/2025/01/TLSI-corporate-deck-011425-Final-2.pdf>
² Internal estimate based current patient data
³ Assuming attaining 33% of RenovoRx estimated addressable market (SAM)

High Margin Business Model

Gross Margins (Actual)	~85%
Required overhead	Low
Operating Margins (Target)	30%

Patented Highly Defensible Position

RenovoRx holds a robust global IP portfolio of 20 issued or allowed patents and 13 pending patents

- Patent protection through 2045
- TAMP-specific cases (covering pressure-mediated trans-arterial delivery) expire 2038
- RenovoRx kit claims expire 2032
- Dual-occlusion device patent expires 2031
- First RenovoRx patent expires December 2030

Disclaimer – We have based our estimates of total addressable market size, peak annual sales projections and similar matters above and elsewhere in this presentation on our market research, third-party reports and publicly available information which we consider reliable. However, readers are cautioned our projected sales and similar metrics are merely our current, preliminary estimates and are subject to many risk factors, many of which are or may be beyond our control. As such, no assurances are given that such estimates will prove to be accurate.

RenovoCath Go-to-Market Strategy

2026 Commercialization Ramp Builds on 2025 Limited Market Launch, Delivering Record Revenue Growth

Concentrated Market: fewer than 200 hospitals treat majority of estimated patients where RenovoCath is seeing interest in utilization



RenovoCath potential high-volume centers¹

Initial target customers have expressed interest in treating a minimum of 6-12 patients with RenovoCath annually (>5 treatments per patient)²

¹ For illustrative purposes only, ² 70+ physician interviews

Go-To-Market Strategy

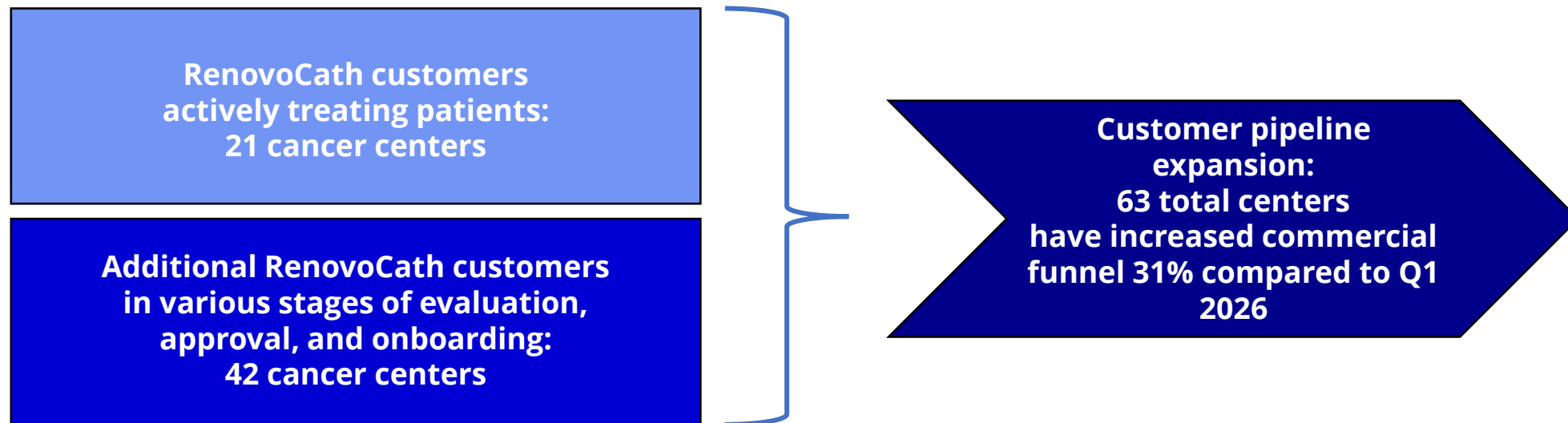
Broadening adoption momentum: Customer pipeline continues to expand across high-volume NCI-designated centers, academic institutions, and large community centers.

Capital-efficient commercialization with disciplined scale-up underway

- **Small/lean sales team (approx. 5 reps)** can tackle top 100+ hospitals
- **Record Q2 2026 revenue of \$909K**, +115% YoY, representing approximately 83% of full-year 2025 revenue
- **1H 2026 revenue reached \$1.5M**, demonstrating strong momentum and continued acceleration
- **Lean, high-performing sales & marketing** team with deep market expertise and a clear growth strategy executed within the existing operating budget to support scalable growth in 2026 and beyond.

RenovoCath Commercialization Update (as of 8/12/26)

2025 Commercial Exploration Laid Foundation for 2026 Revenue Growth and Commercialization



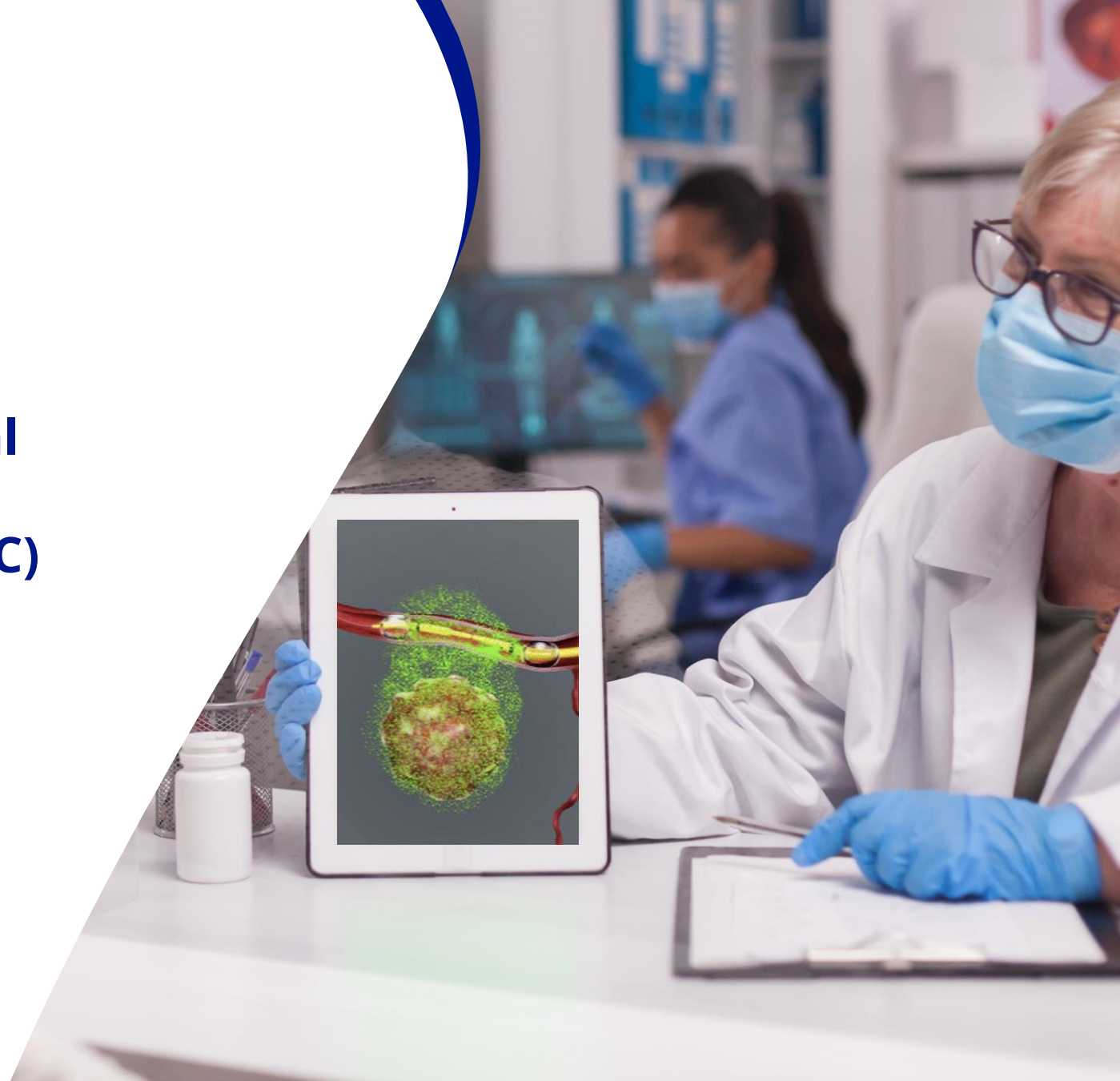
- Key performance indicators predict future revenue: customers utilizing/reordering + approved purchasers + total engaged cancer centers/sales funnel
- Revenue stability & growth: 2026 providing improved visibility into timing of cashflow breakeven, with revenue ramp underway
- With patient enrollment complete, 15 Phase III TIGeR-PaC trial sites that have used RenovoCath are positioned to transition into commercial cancer center customers, several have already begun transitioning
- Increased 2026 annual revenue guidance from a range of \$3M-\$4M to \$3.75M to \$4.25M
- Management believes that reaching a quarterly revenue run-rate of approximately \$5 million would position RenovoRx at cash-flow break-even; based on current trajectory, this could occur in the Q4 2027

Combination Product Pipeline & Investigator Initiated Trial (IIT) Strategy

Centers can bill for the TAMP procedure regardless of the agent administered. To generate new data, RenovoRx is conducting the Phase III TIGeR-PaC trial (fully enrolled August 2026) and supports IITs exploring administration of a variety of agents and tumors.

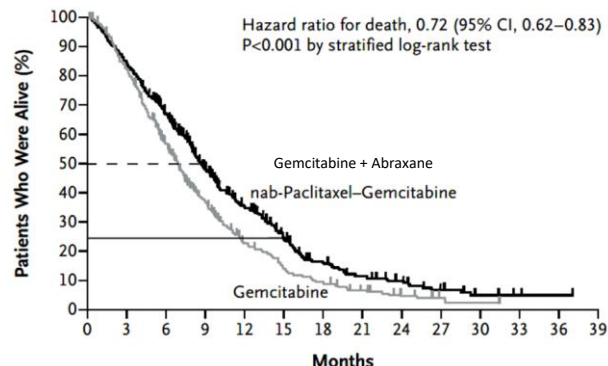
Drug	Indication	Development Stage	Notes
Gemcitabine	Locally Advanced Pancreatic Cancer (LAPC)	Phase III (TIGeR-PaC) (fully enrolled)	Orphan Drug Status
	Borderline Pancreatic Cancer	Two Exploratory IIT collaborations	Orphan Drug Status
	Extrahepatic Cholangiocarcinoma	Exploratory IIT Collaboration	Orphan Drug Status
Oxaliplatin	Metastatic Pancreatic Cancer	Exploratory IIT Collaboration	Orphan Drug Status
Additional drugs being explored: KRAS inhibitors, oncolytic viruses in borderline resectable, locally advanced, and metastatic cancers			

Pivotal Phase III TIGeR-PaC Clinical Trial (NCT03257033) in Locally Advanced Pancreatic Cancer (LAPC)



Significant Unmet Need

Three FDA approvals for first line pancreatic cancer in the last 10 years showed minimal Median Overall Survival (OS) benefit and increased toxicity¹

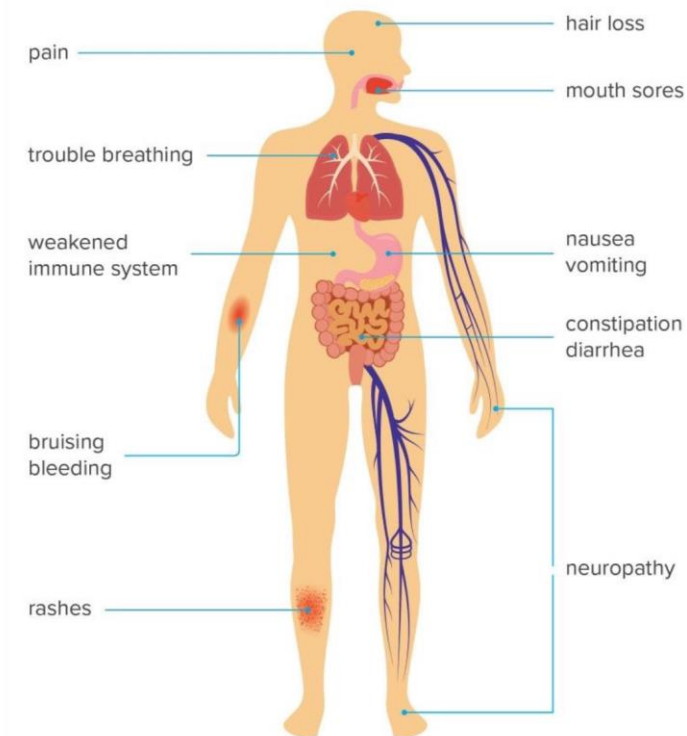


No	Abraxane													
nab-Paclitaxel+Gemcitabine	431	357	269	169	108	67	40	27	16	9	4	1	1	0
Gemcitabine	430	340	220	124	69	40	26	15	7	3	1	0	0	0

Abraxane obtained FDA approval in 2013 on a **7-week** Median Overall Survival (OS) benefit²



38% Grade 3
or Higher Neutropenia and
17% Neuropathy⁴



Olaparib received full FDA approval in 4Q 2019 with no Median OS Difference (<4-mo PFS benefit); Onivyde received FDA approval on a 1.9-mo Median OS benefit in 2015³.

¹ <https://www.cancer.gov/news-events/cancer-currents-blog/2015/irinotecan-liposome-pancreatic>

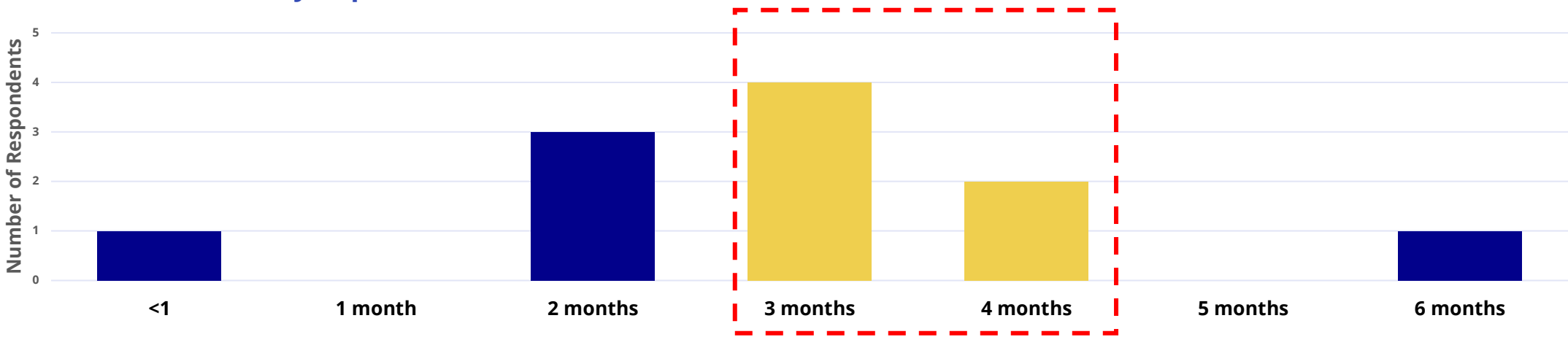
² https://www.accessdata.fda.gov/drugsatfda_docs/nda/2013/021660Orig1s037.pdf

³<https://www.fda.gov/drugs/resources-information-approved-drugs/fda-approves-olaparib-gbrcam-metastatic-pancreatic-adenocarcinoma>

⁴ <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7479547/>

Potential Market Penetration Based on Modest Efficacy from TIGeR-PaC

Independent interviews suggest oncologists likely to adopt gemcitabine delivered via RenovoCath w/ at least 3–4-month OS benefit and toxicity improvement are met



Source – FSI Interviews (multiple responses permitted per respondent)

Testimonials

“Any amount of time, if it is from a phase III. **We started using erlotinib about 15 years ago based on a 14-day benefit.**”

– Abushahin, MedOnc,
The Ohio State University

“The idea of an infusional therapy, **avoiding systemic toxicity is appealing.**”

– Iyer, MedOnc, Roswell

“This targets local vessel involvement and **you can up the delivery of concentration.**”

– Astsaturov, HemOnc,
Fox Chase

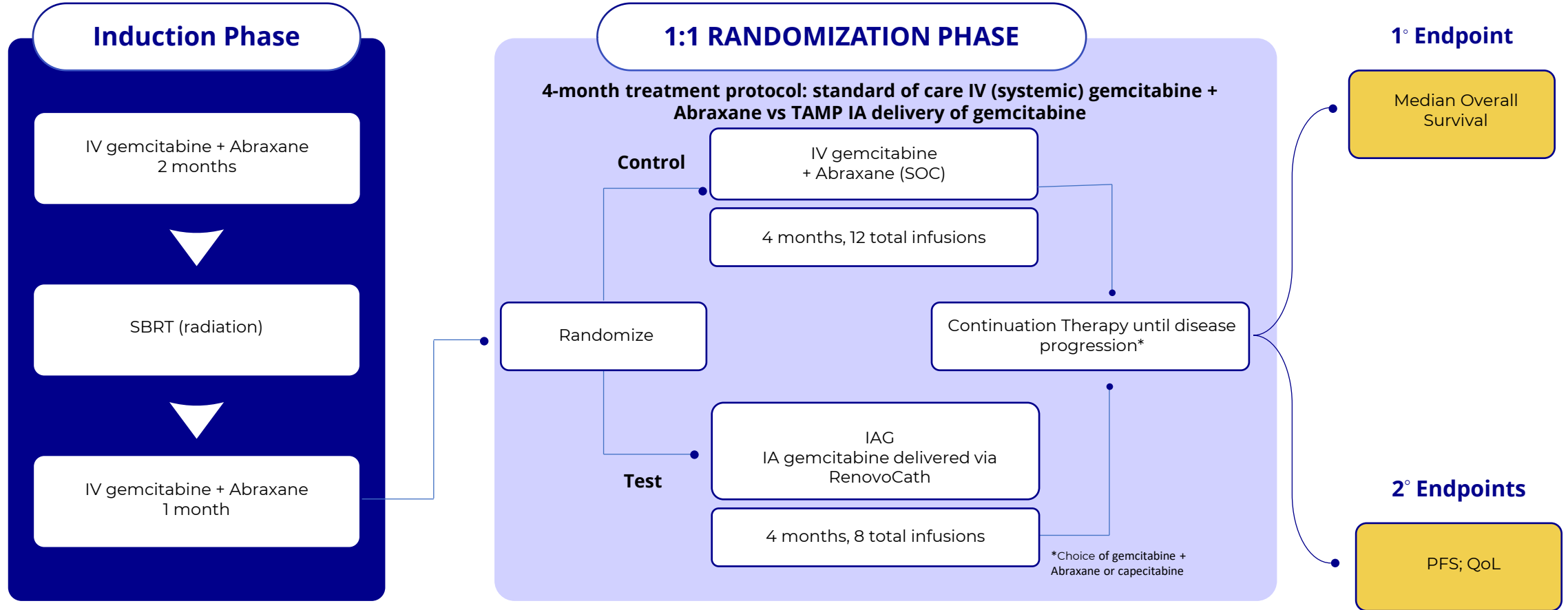
“What do we do with the [unresectable] patients that don’t develop metastatic disease for a while? We’ve essentially talked only about two lines of treatment. But eventually, **toxicity builds up** and they can’t tolerate treatment. For that subset of patients...it would be **highly attractive to offer them something like this.**”

– Mettu, MedOnc, Duke

Disclaimer – This data is based upon independent interviews conducted by Fletcher Spaght, Inc. in 2019. Intra-arterial delivery of gemcitabine via the RenovoCath® is currently in an ongoing Phase III randomized multi-center study evaluating its use for the treatment of locally advanced pancreatic cancer, this drug/device combination has not been approved by the FDA or any regulatory authority and is limited to investigational use only. The mechanism of action for this combination is theoretical only, based on currently available scientific evidence and product design, and may not be reflective of what happens in the human body.

TIGeR-PaC Phase III Randomized Trial Design

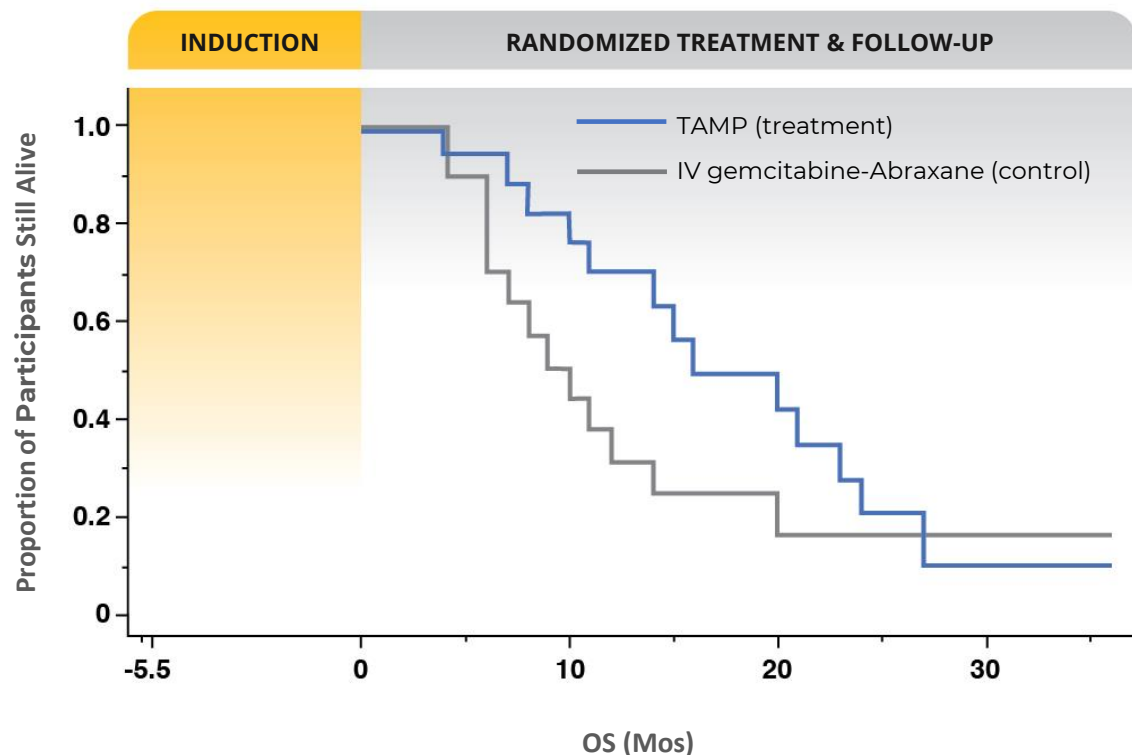
Combination Product Trial for TAMP Delivery of Intra-Arterial (IA) Gemcitabine (IAG) via FDA-Cleared RenovoCath in Locally Advanced Pancreatic Cancer (LAPC)



First Pre-Planned Interim Analysis presented at AACR and ESMO GI 2023

TAMP (IA Gemcitabine via RenovoCath): 6-Month Median OS Benefit vs. IV (Systemic) Gemcitabine + Abraxane

TIGeR-PaC phase III first interim analysis



Data Presented at AACR 2023 and ESMO GI 2023

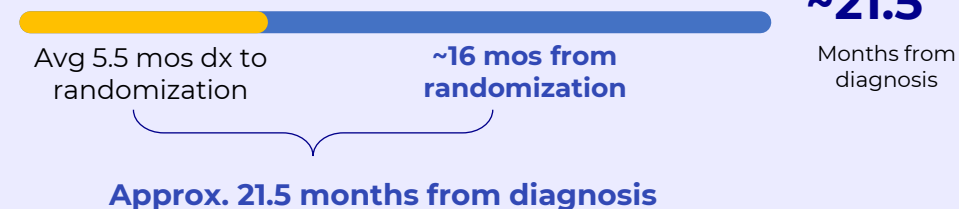


Data on 45 patients randomized

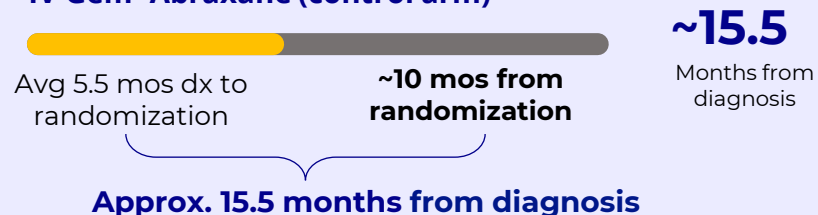
- 23 randomized to RenovoCath plus Gemcitabine
- 22 randomized to IV gemcitabine + Abraxane

Median Overall Survival Difference: 6 Months

TAMP (test arm)



IV Gem+Abraxane (control arm)

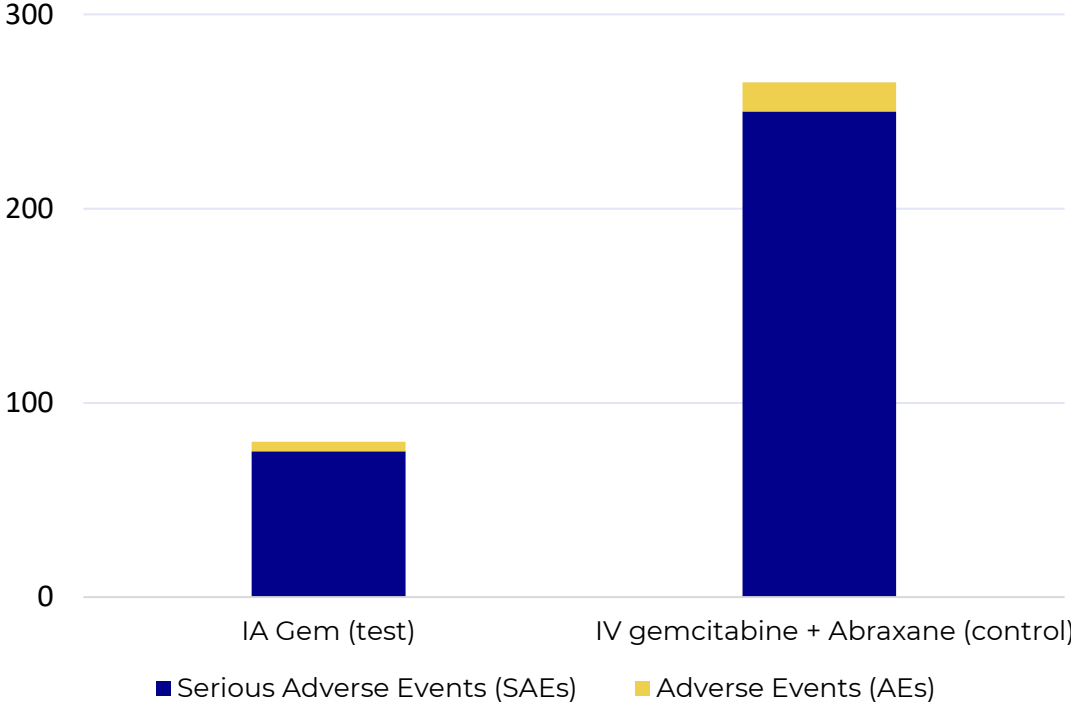


Statistical significance was not reached to stop the study early

TAMP (IA Gem) via RenovoCath: Fewer AEs and SAEs vs IV (systemic) Gemcitabine + Abraxane

TIGeR-PaC phase III first interim analysis

65% fewer total AEs and SAEs in IA vs. IV arm



Data Presented at AACR 2023 and ESMO GI 2023

Fewer AEs in 11/13 categories with greater than 10% frequency in each arm (All Grades)

Adverse Events	IV Gem + Pac	IA Gem
Neutropenia	81%	21%
Anemia	48%	8%
Thrombocytopenia	38%	4%
Elevated AST	33%	4%
Elevated ALT	29%	13%
Fatigue	19%	8%
Neuropathy	19%	0%
Dehydration	19%	8%
Hypertension	14%	4%
Hypokalemia	14%	4%
Hypoalbuminemia	14%	4%
Abdominal Pain	0%	21%
Nausea	10%	17%

Post-Marketing Registry Study (RR5) & Support of Investigator-Initiated Trials (IITs)

Evidence-Based Growth Strategy

Objective

- Expand real-world safety and efficacy data of TAMP in patients diagnosed with solid tumors

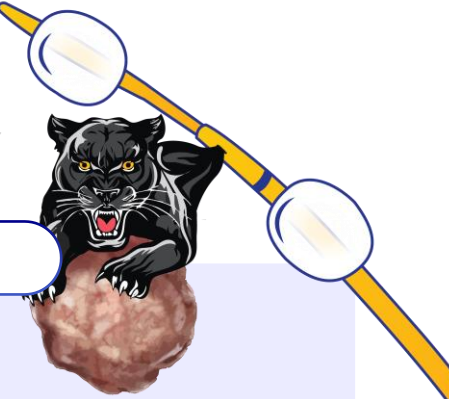
Commercialization Efforts

- Could enable label expansion
- Inform future trials
- Support reimbursement and broader adoption

Clinical and Strategic Advancement

Participating Clinical Sites

- University of Vermont Cancer Center – borderline pancreatic cancer
- Baptist Health Miami Cancer Institute – solid tumor registry
- University of Pittsburgh Medical Center – solid tumor registry
- Moffit Cancer Center, Tampa – metastatic pancreatic cancer
- City of Hope Cancer Center – cholangiocarcinoma (bile duct cancer)



"This study provides a crucial opportunity to evaluate how RenovoCath can improve drug-delivery in patients diagnosed with solid tumors, while potentially, and importantly, improving survival and quality-of-life outcomes."

–Dr. Conor O'Neill of the University of Vermont Cancer Center

**Capital-efficient studies providing meaningful data that may further broaden the application
for the TAMP therapy platform enabled by RenovoCath**

RNXT Financial Highlights

Record Q2 2026 Revenue of \$909K, up 115% Year-Over-Year

- Driven by accelerating commercial adoption
- 1H 2026 revenue: \$1.5 million, up 138% over 1H 2025;
Q2 2026 revenue equaled approximately 83% of full-year 2025 revenue
- Phase III TIGeR-PaC trial reaches full enrollment
 - Completion of trial expected in first half 2027 with topline data readout expected in second half 2027

Executed On All 3 Milestones

- ✓ **Revenue Growth:** Delivered another record revenue quarter
- ✓ **Commercial Momentum:** Achieved several new cancer center activations
- ✓ **Technology Expansion:** Expanded the application of our technology beyond locally advanced pancreatic cancer (LAPC)

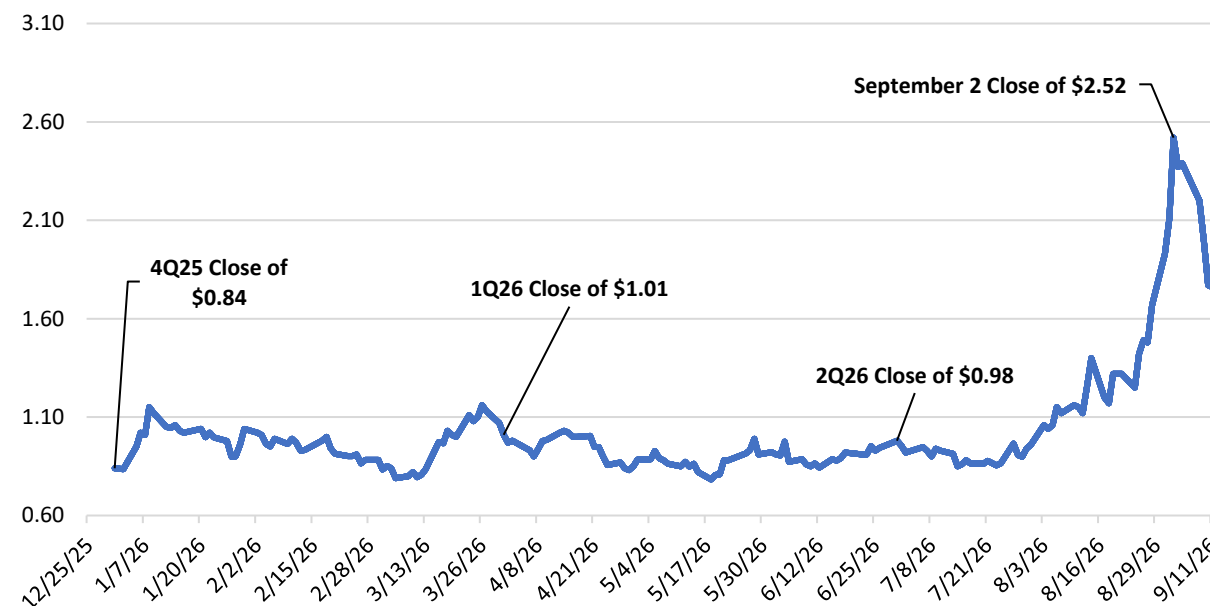
Ownership

- Institutional: 20.4% or ~9.2M shares¹
 - Backed by multiple fundamental healthcare institutional investors
- Insider: 3.61% or ~1.6M shares¹

Increased 2026 Outlook

- Increased revenue guidance from \$3-4M to \$3.75-4.25M
- Strong commercial activity: Targeting at least 36 active commercial cancer centers by year-end; footprint of 21 active commercial cancer centers as of August 12, 2026

RNXT (USD) Daily Performance - 2026 YTD through September 11, 2026



New 52-Week High:	\$2.59
Average Daily Volume Since Q2 Results on 8/12:	1.44M shares
Market Cap:	\$87.1M ¹
Shares Outstanding:	45.13M ¹

¹ As of 8/31/26

Experienced Management Team



Shaun R. Bagai

Chief Executive Officer & Board Member

- HeartFlow (>\$2B IPO)
- Ardian (Acquired for > \$900M)
- Medtronic Vascular
- TransVascular (Acquired by Medtronic)



Ramtin Agah, MD

Chief Medical Officer, Founder & Executive Chairman

- Interventional Cardiology, El Camino Hospital
- Consultant, Abbott Vascular



Mark Voll

Chief Financial Officer

- Techwell, Inc. (IPO in 2006, acquired by Intersil in 2010)
- Montage Technology (IPO in 2013, acquired by PDSTI in 2014)
- Aquantia Corporation (IPO in 2017, acquired by Marvell Technology in 2019)



Leesa Gentry

Chief Clinical Officer

- Evotec
- Otsuka America Pharmaceuticals
- Omnicare Clinical Research



Richard Stark

Commercial Advisor Consultant

- Innobative Designs
- AngioDynamics



Ronald B. Kocak, CPA

Vice President, Controller & Principal Financial Officer

- Sensei Biotherapeutics, Inc.
- Member of the American Institute of Certified Public Accountants
- Member of Chartered Global Management Accountant



Thomas Kelly

Vice President, Quality & Operations

- Perclose/Abbott Vascular, Cierra, Arstasis, Tioga Cardiovascular
- Stryker (Orthopedics)
- ForSight Vision4/Genentech (Ophthalmology)
- AEGEA/CooperSurgical (Women's Health)



Phil Stocton

Vice President, Sales & Market Development

- Sirtex
- Varian
- AngioDynamics
- Nipro Corporation

Board of Directors



Ramtin Agah, MD

Executive Chairman, Chief Medical Officer, & Founder

- Interventional Cardiology, El Camino Hospital
- Consultant, Abbott Vascular



Laurence J. Marton, MD

Director

- Board: Cellsonics, TOMA Biosciences, xCures



Shaun R. Bagai

Director & Chief Executive Officer

- HeartFlow (>\$2B IPO)
- Ardian (Acquired for > \$900M)
- Medtronic Vascular
- TransVascular (Acquired by Medtronic)



Una S. Ryan, PhD, OBE

Director

- Board: Cortexyme, Elemental Machines



Angela Macfarlane

Director

- CEO, Perceive Biotherapeutics (\$78M raised led by JJDC)
- CEO, Foresight Labs



Robert J. Spiegel, MD

Director

- CMO, PTC Therapeutics
- CMO, Schering-Plough (\$41.1B merger with Merck)

RenovoRx Scientific Advisory Board



Timothy Donahue, MD

- Director of the UCLA Agi Hirshberg Center for Pancreatic Diseases
- Garry Shandling Chair in Pancreatic Surgery, David Geffen School of Medicine at UCLA



Karyn A. Goodman, MD, MS

- Professor and Vice Chair of Clinical Research, Department of Radiation Oncology, Icahn School of Medicine at Mount Sinai
- Associate Director of Clinical Research, The Tisch Cancer Institute at Mount Sinai



Michel Ducreux, MD, PhD

- Head of the Gastrointestinal Oncology Unit and Gastrointestinal Oncology Tumor Board at Gustave Roussy
- Professor of Oncology at Paris-Saclay University in France
- Vice-Chair of ESMO GI



Mike Pishvaian, MD, PhD

- Associate Professor, Department of Oncology Director of the Gastrointestinal, Developmental Therapeutics, and Clinical Research Programs at the NCR Kimmel Cancer Center at Sibley Memorial Hospital
- Johns Hopkins University School of Medicine



Margaret A. Tempero, MD

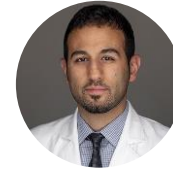
- Professor of Medicine and Director of the UCSF Pancreas Center
- Editor-in-Chief of JNCCN
- Former ASCO President

RenovoCath Medical Advisory Board



Nadine Abi-Jaoudeh, MD

- Chief of Interventional Radiology and Director of Clinical Research in the Department of Radiology at the University of California, Irvine



Mustafa Al-Roubaie, MD

- Interventional Radiologist at Moffitt Cancer Center
- Associate Professor of Radiology and Oncologic Sciences at the University of South Florida School of Medicine



Khashayar Farsad, MD, PhD

- Josef Rosch Professor and Chair of the Department of Interventional Radiology, Oregon Health and Science University
- Director of Dotter Interventional Institute, and Director of Research, Oregon Health and Science University



Ripal Gandhi, MD

- Interventional Radiologist at Baptist Health South Florida



Paula Marie Novelli, MD

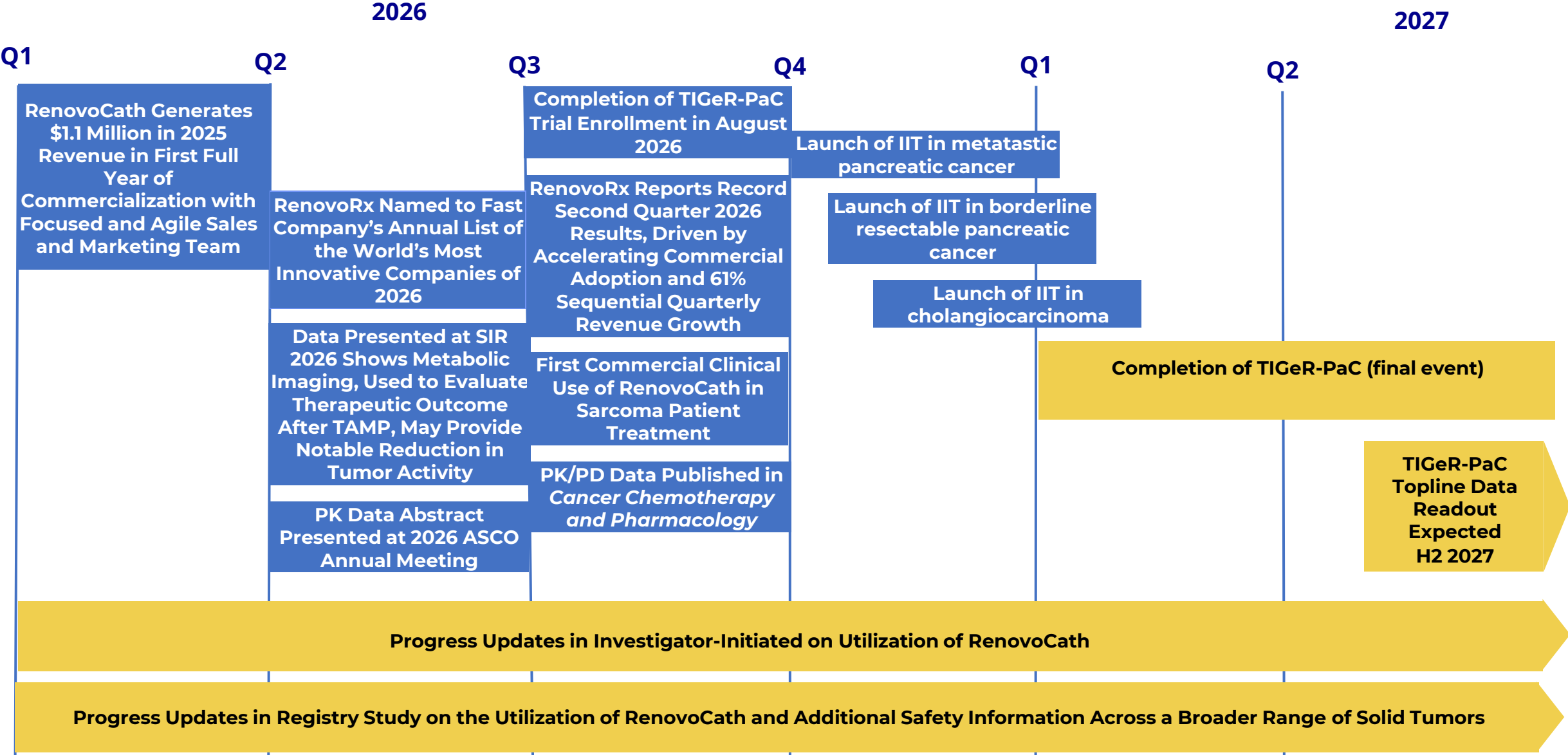
- Interventional Radiologist at University of Pittsburgh Medical Center



Jonathan Kessler, MD

- Chief of the Division of Interventional Radiology at City of Hope Comprehensive Cancer Center

RNXT 2026-2027 Milestones



How Hospitals are Using TAMP with RenovoCath in Oncology Management*

Visit <https://ir.renovorx.com/news-events/clinical-news> to Learn More



NBC News highlighted the potential of RenovoRx's Trans-Arterial Micro-Perfusion (TAMP) therapy platform, designed for targeted drug-delivery via the RenovoCath device in difficult-to-treat tumors, like pancreatic cancer. Dr. Gregory Tiesi, Medical Director of Hepatobiliary Surgery, is interviewed about his patient receiving treatment with TAMP at Hackensack Meridian Health.



Spectrum News 1 SoCal recently featured an interview with Dr. Jonathan Kessler, Chief of Interventional Radiology at City of Hope, highlighting how TAMP therapy platform, enabled by RenovoCath, has the potential to reshape oncology management for patients diagnosed with difficult-to-treat tumors, like pancreatic cancer.



"Trans-Arterial Micro-Perfusion is a drug-delivery platform that delivers chemotherapy directly near the target tumor." Dr. Ripal Gandhi, a vascular interventional radiologist at Baptist Health Miami Cardiac & Vascular Institute and Miami Cancer Institute, spoke with Fox News about the challenges of treating pancreatic cancer and the potential of the TAMP platform as a targeted drug-delivery option.

The RenovoCath® device is indicated for temporary vessel occlusion in applications including arteriography, preoperative occlusion, and chemotherapeutic drug infusion. RenovoRx is evaluating the use of RenovoCath-enabled TAMP with gemcitabine as a combination product (known as intra-arterial gemcitabine) for the treatment of locally advanced pancreatic cancer (LAPC), including in the ongoing Phase III TiGeR-PaC trial, under a U.S. Investigational New Drug Application. Ongoing clinical studies continue to evaluate safety and clinical outcomes in patients with LAPC. For the full FDA-cleared indications for use for RenovoCath and associated contraindications, warnings, and precautions please visit: <https://renovorx.com/#ifu>



Thank You

Investor Contact:

KCSA Strategic Communications

Valter Pinto or Jack Perkins | RenovoRx@KCSA.com

+1.212.896.1254

Media Contact:

StiR Communications

Hannah Williams | Hannah@stir-communications.com

+1.803.521.1214