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We've made improving lives our responsibility.
And that drives our passion and success

A mission, a plan, and crisp execution producing sustained growth



- ✓ 6 Large & Underpenetrated Markets
- ✓ Purpose-Built& Highly Differentiated Toolkits
- **✓** Robust Pipeline of Innovation**

- ✓ Compelling Clinical Evidence
 7 Leading Clinical Studies***
- ✓ Significant Commercial Growth 20+ International Markets
- √ 1,300+ Employees and Growing

^{*}Based on third party data and Inari management estimates

^{**} Products launched 2021: Triever20 Curve catheter, FlowTriever2 catheter, FlowStasis, FlowSaver, Triever24 Flex catheter; 2022: ClotTriever BOLD catheter, Intri24 sheath, Protrieve sheath, InThrill system, Triever Gen 4 catheters; 2023:Triever 16 Curve catheter, RevCore, ClotTriever XL catheter, ClotTriever BOLD Gen 2 catheter

^{***} FLARE, FLASH, FLAME, CLOUT, PEERLESS RCT, PEERLESS II RCT, DEFIANCE RCT

Venous Thromboembolism (VTE)

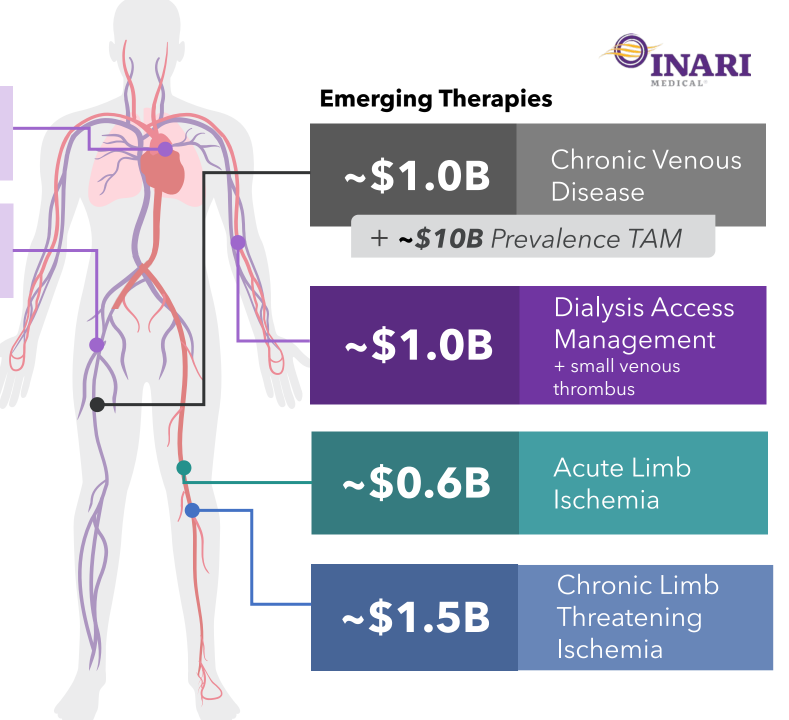
~\$2.8B

Pulmonary Embolism

~\$3.0B

Deep Vein Thrombosis

~\$10B U.S. TAM across 6 disease states





Three growth pillars supported by global commercial playbook

1

Venous
Thromboembolism (VTE)

Driving our solutions towards standard of care in VTE

- Pulmonary Embolism (PE)
- Deep Vein Thrombosis (DVT)

2

Emerging Therapies

Building momentum with new products in new markets

- Chronic Venous Disease (CVD)
- Dialysis Access Management (DAM)
- Acute Limb Ischemia (ALI)
- Chronic Limb Threatening Ischemia (CLTI)

3

International

Expanding our footprint internationally

- EMEA: ~20 active countries
- APAC & LAC: ~12 active countries
- China & Japan commercial in 2024

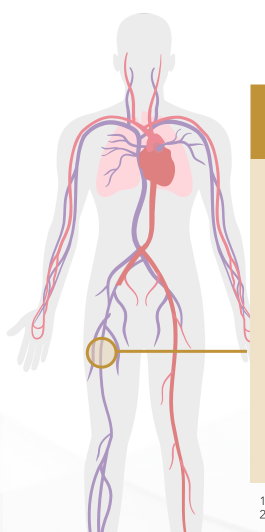
Commercial Playbook

- Expanding our commercial footprint
- Standardizing patient pathways
- Executing guideline changing clinical trials
- Developing purpose-built solutions

Venous Thromboembolism (VTE)

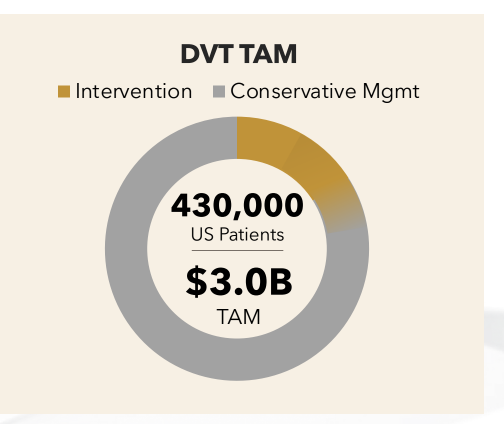
Transforming the lives of patients suffering from DVT





DEEP VEIN THROMBOSIS (DVT)

- A/C alone leaves clot behind in up to half of patients¹
- Lytics don't address chronic clot, and come with bleeding risk
- Up to 50% develop Post-Thrombotic Syndrome (PTS)²

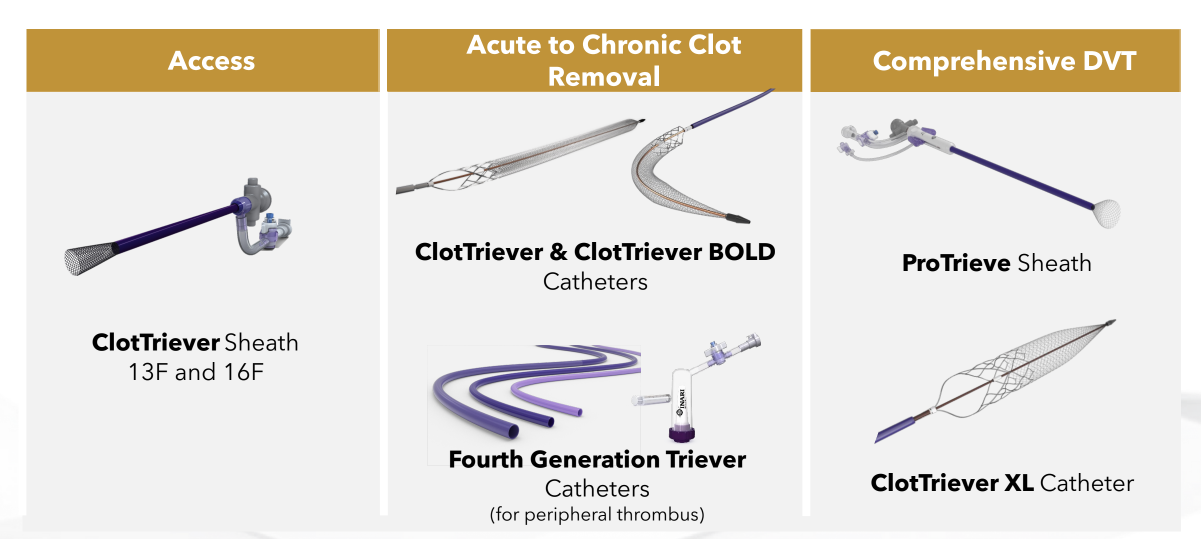


^{1.} Young et al., Post-treatment residual thrombus increases the risk of recurrent deep vein thrombosis and mortality. J Thromb Haemost 2006; 4: 1919-24.

^{2.} Kahn, Susan R. Hematology Am Soc Hematol Educ Program. 2016 Dec 2; 2016(1): 413-418

The ClotTriever® System: A complete solution for DVT and peripheral venous thrombus





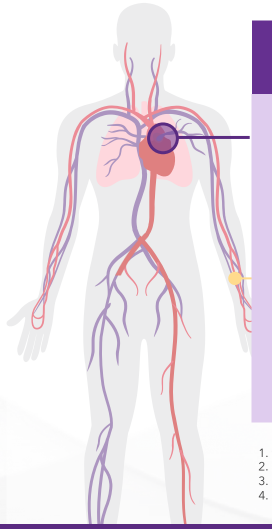
INDICATIONS FOR USE: The ClotTriever thrombectomy system is indicated for: (1) The non-surgical removal of thrombi and emboli from blood vessels. (2) Injection, infusion, and/or aspiration of contrast media and other fluids into or from a blood vessel. The Protrieve Sheath is indicated for use is a conduit for the insertion of endovascular devices into the vasculature while minimizing blood loss associated with such insertions. The FlowTriever retrieval/aspiration system is indicated for (1) the peripheral vasculature moves of emboli and thrombi from blood vessels (2) injection, infusion and/or aspiration of contrast media and other fluids into or from a blood vessel. The FlowTriever retrieval/aspiration system is intended for use in the peripheral vasculature and for the treatment of pulmonary embolism. Triever catheters are also intended for use in treating clot in transit in the right atrium, but not in conjunction with FlowTriever actheters. The FlowTriever above the respective owners.

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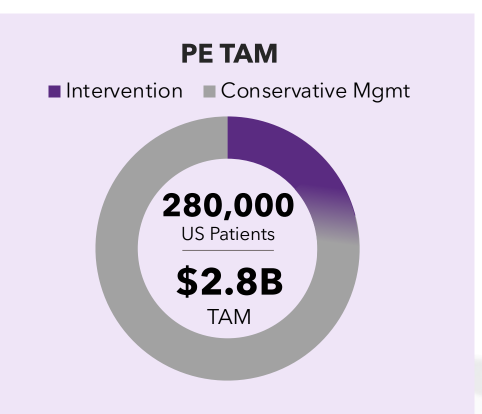
Transforming the lives of patients suffering from PE





PULMONARY EMBOLISM (PE)

- 3rd leading cause of cardiovascular death¹
- A/C alone leaves clot behind in up to half of patients^{2,3}
- Long-term complications are common⁴

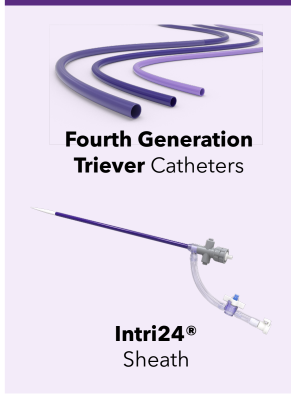


- 1. "Pulmonary Embolism in 2017: Increasing Options for Increasing Incidence", National Center for Biotechnology Information, May 2017.
- 2. Picart, et al. Predictors of residual pulmonary vascular obstruction after pulmonary embolism: Results from a prospective cohort study. Thrombosis Research. 2020.
- 3. Dzikowska-Diduch, et al. The post-pulmonary syndrome results of echocardiographic driven follow up after acute pulmonary embolism. Thrombosis Research. 2020.
- 4. Sista AK, et al. Vasc Med. 2017 Feb;22(1):37-43

The FlowTriever® System: A full toolkit approach to PE







Large Clot Hauls Without Lytics



Large Bore Aspiration



Large Bore Syringe and **Whoosh** Mechanism

Address Challenging Clot or Anatomy



Triever20 Curve® and Triever16 Curve®

Catheters



FlowTriever
Catheters

Minimal Blood Loss



FlowSaver® Blood Return System



FlowStasis® Suture Retention Device

We remove the full range of clot chronicity



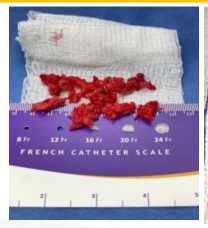
Acute Chronic

ClotTriever® System











FlowTriever® System









High impact clinical evidence to change standard of care













DVT STUDIES

Largest
Prospective PE
Device Study

Largest
Prospective
High-risk PE
Device Study

First & Only Headto-Head Advanced Therapy RCT (FlowTriever v. CDT) RCT Designed to Establish Standard of Care

(FlowTriever v. AC alone)

Largest
Prospective DVT
Thrombectomy
Study

First Industry
Sponsored DVT
RCT
(ClotTriever v. AC)

~1,000 Patients **~80** Sites

115 Patients11 Sites

550+ Patients **60** Sites

1,200 Patients Up to **100** Sites

500 Patients **47** Sites

300 Patients **60** Sites

Enrollment Complete

Enrollment Complete

Enrollment Complete

Enrolling

Enrollment Complete

Enrolling

~3,800 patients across 6 studies

VTE Excellence is a codified & scalable





ENGAGE

process

~1,400 Inari Accounts

Find champions, build the foundation

Average
Account TAM
Penetration

EMPOWER

~250 Inari Accounts

Create patient pathway and build awareness



EXCEL

~50 Inari Accounts

Solidify consistent patient identification, triage, tracking



Average Account TAM Penetration

Our VTE solutions confer significant benefits to hospitals, physicians, and patients



Safely capture and remove large clot burden



Effective, short, single-session treatments



Thrombolytic-free treatment approach



Avoid lytic-based ICU stay



Established procedural reimbursement

Our products offer benefits and value to our hospital and physician customers



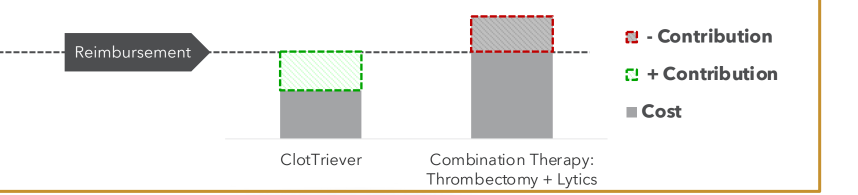
Established Coding & Payment for Mechanical Thrombectomy*

Total Cost/Reimbursement Comparison
Illustrative Procedural Hospital Contributions*

DVT Payment

\$17,080 - \$35,406

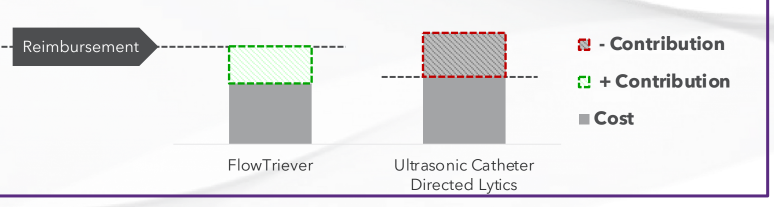
DRG: 270 - 272



PE Payment

\$13,138 - \$33,003

DRG: 163 - 165



^{*} Utilizes national average Medicare reimbursement rates FY2024 IPPS FR and Inari management estimates around patients with and without MCC and CC.

Treatment of thrombotic diseases consistently evolves to definitive catheter based intervention



Anticoagulation (AC) Only

AC +
Thrombolytics (Lytics)

AC + **Definitive Catheter Intervention**



Myocardial Infarction

AC alone

AC + Thrombolysis

AC + **POBA & DES**





Stroke

AC alone

AC + **Systemic Lytics**

AC + Lytics +
Stentrievers & Aspiration
Thrombectomy





AC alone

AC +
Systemic & Catheterdirected Lytics

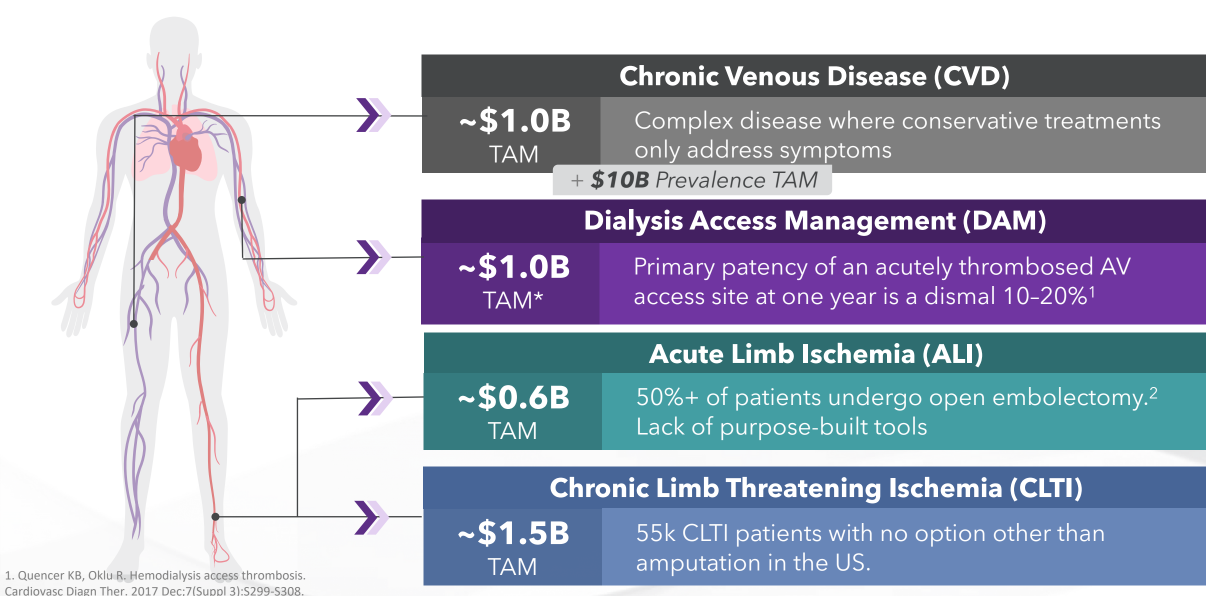
AC +
ClotTriever &
FlowTriever



Emerging Therapies

Emerging Therapies

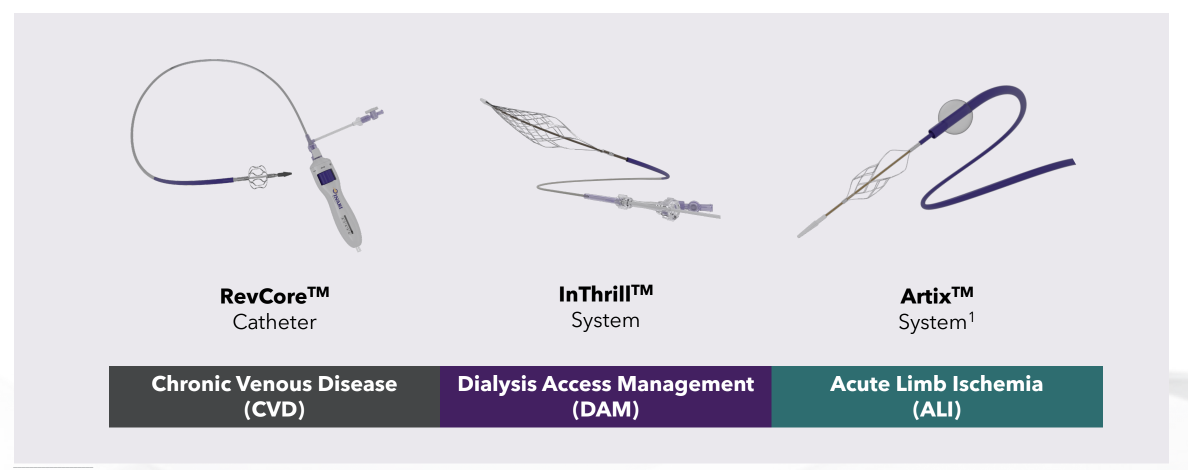




^{2.} Based on third party data and Inari management estimates.



Organically expanding beyond VTE



¹⁾ Artix is not currently being marketed, and is expected to launch in 2024.

INDICATIONS FOR USE: The RevCore Thrombectomy Catheter is indicated for: (1)The non-surgical removal of thrombi and emboli from blood vessels. (2) Injection, infusion, and/or aspiration of contrast media and other fluids into or from a blood vessel. The RevCore Thrombectomy Catheter is indicated for (1) the non-surgical removal of thrombi and emboli from blood vessels, including arteriovenous fistulae and arteriovenous grafts for dialysis access, and synthetic grafts and (2) injection, infusion, and/or aspiration of contrast media and other fluids into or from a blood vessel, graft. The InThrill Thrombectomy System is intended for use in the peripheral vasculature. The InThrill Thrombectomy device is indicated for (1) the non-surgical removal of emboli and thrombi from peripheral vasculature. The InThrill Thrombectomy device is indicated for (1) the non-surgical removal of emboli and thrombi from peripheral vasculature. The InThrill Thrombectomy device is indicated for (1) the non-surgical removal of emboli and thrombi from peripheral vasculature. The InThrill Thrombectomy device is indicated for (2) injection, infusion, and/or aspiration of contrast media and other fluids into or from a blood vessel, the non-surgical removal of emboli and thrombi from peripheral vasculature. The InThrill Thrombectomy device is indicated for use in deplication of contrast media and other fluids into or from a blood vessel, the non-surgical removal of emboli and thrombic from peripheral vasculature. The InThrill Thrombectomy device is indicated for use in the peripheral vasculature. The InThrill Thrombectomy device is indicated for use in the peripheral vasculature. The Artix BG balloon guiding sheath is indicated for use in the peripheral vasculature. The Artix BG balloon guiding sheath is indicated for use in the peripheral vasculature. See Instructions for Use for complete Indications, warnings, and precautions. Caution: Federal (USA) law restricts this device to sale by or on the order of a physician. All trademarks are prop

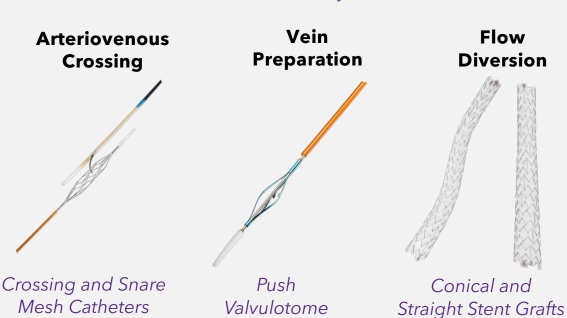
LimFlow System - Transforming the Treatment of CLTI



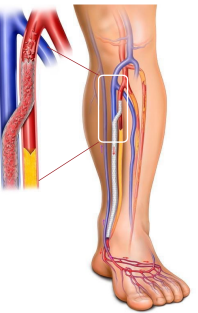
Chronic Limb Threatening Ischemia (CLTI): The LimFlow® System

Transcatheter Arterialization of Deep Veins (TADV)

with the LimFlow System:



Arterialized Veins Post-LimFlow

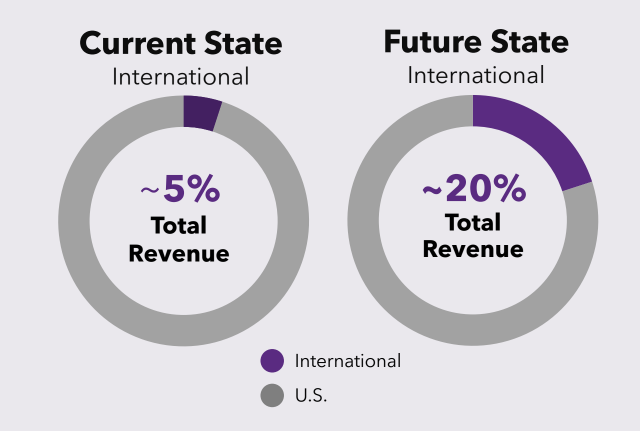


LimFlow System Highlights

- Call Point: Vascular surgery & interventional radiology / cardiology
- Site of Service: Primarily hospital-based peripheral interventions
- Only On-Label Device for No-Option CLTI. FDA PMA Approved in Sept. 2023
- PROMISE II study published in NEJM, the world's leading medical journal

International Markets

A vast global unmet need offers a significant runway for growth



Key Drivers:

Level 1 RCT data

Changing Guidelines

Incremental Reimbursement in Key Geographies

Entering Remaining Key Markets: China and Japan



Q4 2023 Financial Update







2023 **Q4 Revenue**

\$132.1M

~23% increase over 2022 Q4

2023 FY Revenue

\$493.6M

~29% increase over full year 2022





2024 FY Revenue Guidance

\$580M - \$595M

17.5% - 20.5% increase over full year 2023

Financial Profile

- Exceptional growth, significant runway
- Premium 85%+ gross margin profile
- Solid core cash flow generation to support LimFlow and growth objectives
- Sustained operating profitability by 1H 2025

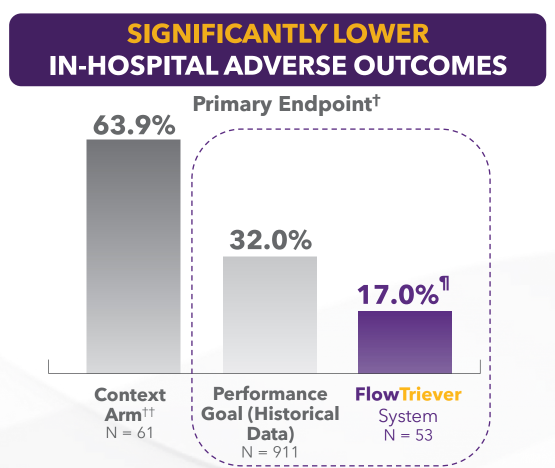


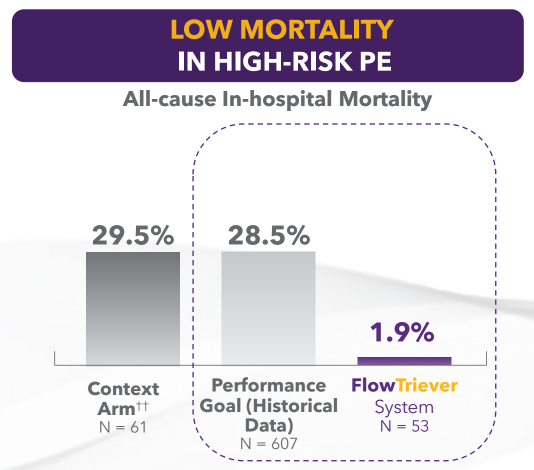


FLAME high-risk PE study shows very low rates of adverse events and mortality



Results from FLAME: The largest prospective study of interventional treatment in high-risk PE





Composite primary endpoint consisted of in-hospital all-cause mortality, bailout to an alternate throubus removal strategy, clinical deterioration, and major bleeding

Context arm patients were treated with systemic thrombolysis (68.9%), anticoagulation alone (23.0%), CDT (6.6%) or surgical thrombectomy (1.6%) >20.01 vs. performance gnal hased on historical data

Source: Silver, M J et al., Outcomes in High-risk Pulmonary Embolism Patients Undergoing FlowTriever Mechanical Thrombectomy or Other Contemporary Therapies: Results from the FLAME Study. Circ. Cardiovasc. Interv. 2023 O

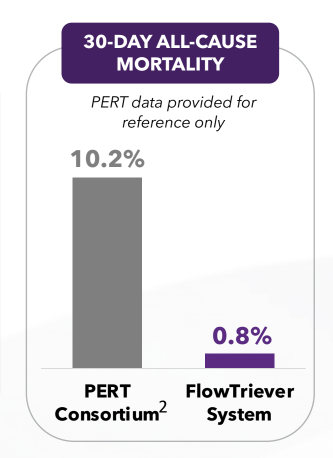
FLASH is the largest prospective registry in PE with exceptional results¹

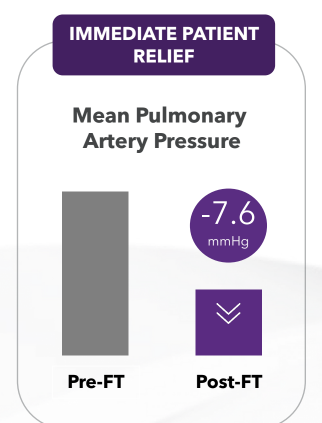


800 patients, 50 sites, 32% were contraindicated to lytics³



0%
Device related MAEs







90%
Mild or absent dyspnea at 6 months

1.0%

CTEPH at 6 months

^{1.} Toma C, et al. Acute Outcomes for the Full US Cohort of the FLASH Mechanical Thrombectomy Registry in Pulmonary Embolism. EuroIntervention 2023;18:1201-1212.

^{2.} PERT Consortium Quality Database. October 2021 (Presented by Secemsky E); Darki A & Jaber WA. Endovascular Today. July 2022 Supplement (PERT Updates)

^{3.} Represents number of patients in the full US cohort.

CLOUT is the largest mechanical thrombectomy dataset in DVT with exceptional results¹



500 patients | **47** sites | **70%** subacute and/or chronic clot | **30%** lytics contraindicated

EXCELLENT SAFETY RESULTS

0.2%

Device related SAEs

0% valve damage0% vessel damage0% acute kidney injury

0.4% Thrombolytics used

EXTENSIVE CLOT REMOVAL, REGARLESS OF CLOT AGE

Overall

>90%

Complete or Near Complete Thrombus Removal* By clot age**

91% in acute

82% in subacute

84% in chronic

* ≥75% thrombus removal

EXCELLENT OUTCOMES†

>90%

Freedom from moderate or severe PTS symptoms through 1 year (N=227)

95% Flow via duplex ultrasound at 1 year

N = 192

^{1.}Dexter D, Kado H, Shaikh A, et. al., Safety and Effectiveness of Mechanical Thrombectomy From the Fully Enrolled Multicenter, Prospective CLOUT Registry Journal of the Society for Cardiovascular Angiography & Interventions, Volume 2, Issue 2, March-April 2023, 100585

^{**}Subset of 250 patients presented at AVF 2022

[†] One-year interim outcomes from the multicenter prospective CLOUT registry presented by Dr. David Dexter VEINS 2023



Superiority RCT of FlowTriever vs CDT in PE

550 Patients Randomized 1:1

FlowTriever

CDT

Patients Followed for 30 Days

HIGHLIGHTS



Currently, Catheter Directed Thrombolysis (CDT) is used in ~40% of interventions commercially*



Primary endpoint via win ratio:

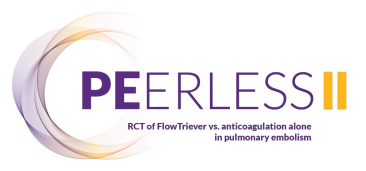
- All-cause mortality
- Intracranial hemorrhage
- ISTH major bleeding
- Clinical deterioration/bailout
- ICU admission & ICU LOS



Enrollment nearly complete



Designed to transform standard of care away from CDT



RCT of FlowTriever vs Anticoagulation Only in PE

Intermediate-risk Acute PE

1,200 Patients Randomized 1:1

FlowTriever

AC

Patients followed for 3 months

HIGHLIGHTS



Currently, anticoagulation alone is the guideline-recommended therapy for intermediate-risk PE patients



Primary endpoint via win ratio:

- All-cause mortality
- Clinical deterioration
- All-cause hospital readmission
- Bailout therapy
- Dyspnea score



Enrollment starting soon



Designed to transform standard of care



Superiority RCT of ClotTriever vs Anticoagulation in DVT

Moderate-Severe Iliofemoral DVT

300 Patients Randomized 1:1

ClotTriever

Conservative Medical Management (A/C)

Patients Followed for 6 Months

HIGHLIGHTS



First global industrysponsored RCT for DVT



Primary endpoint via win ratio:

- Treatment failure or escalation of therapy
- Post-Thrombotic Syndrome severity at 6 months



Enrollment started in January 2023



Designed to transform standard of care

PROMISE II: Pivotal Study at 6 Months

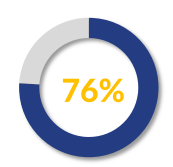


- ➤ Landmark multi-center, prospective pivotal trial of the LimFlow System conducted at 20 U.S. centers in 105 No-option CLTI patients typically excluded from other clinical studies. ¹
- All patients were confirmed as "No-Option" and facing imminent amputation by an independent review committee of vascular surgeons. 1
- 6 Month results published in the New England Journal of Medicine

Limb Salvage at 6 Months Wou

Functional Limb Preservation in No-Option Patients

Wounds Healed or Healing at 6 Months



Wound Healing in Patients
With Non-Healing Chronic
Wounds

Technical Success



No Device-related Adverse Events

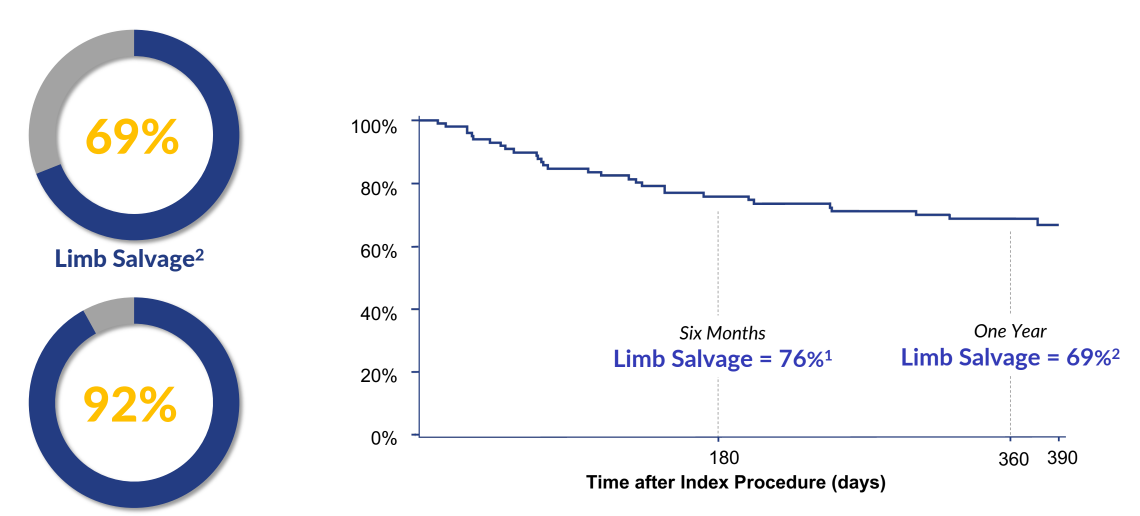
Sickest population of CLTI patients ever enrolled in a pivotal trial.



2023

PROMISE II: Durability of Limb Salvage Results Demonstrated at 12 Months





Wounds Healed or Healing²