

# EKOS™ Endovascular System

## KNOCOUT PE

### International EkoSonic Registry of the Treatment and Clinical Outcomes of Patients with Pulmonary Embolism

#### Prospective Cohort 3-month Data Release

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#### TRIAL OBJECTIVE:

To understand the impact of the OPTALYSE PE study on various ultrasound-accelerated thrombolysis (USAT) protocols being used as the standard of care in the treatment of acute PE and associated long-term outcomes.

#### CENTERS:

83 international sites across the United States and Europe

#### PATIENTS:

Retrospective cohort: **991 patients**  
Prospective cohort: **489 patients**

#### REGISTRY DESIGN | PROSPECTIVE COHORT

Patients	Inclusion Criteria	Exclusion Criteria
<p><b>489 patients</b> with intermediate-high risk and high-risk PE</p> <p>Treated with EKOS from March 2018-June 2020</p>	<ul style="list-style-type: none"> <li>• Male or female &gt; 18 years of age and &lt; 80 years of age</li> <li>• Intermediate High-Risk or High-Risk PE</li> <li>• RV/LV &gt; 1.0 from diagnostic CTA or echocardiogram</li> <li>• Symptom duration &lt; 14 days</li> <li>• Troponin elevation</li> <li>• Investigator has selected the EKOS device to treat patient</li> <li>• Infusion dose/duration per investigator's SOC</li> </ul>	<ul style="list-style-type: none"> <li>• Clinician deems subject high-risk for catastrophic bleeding</li> <li>• Life expectancy &lt; 1y</li> </ul>

#### END POINTS

Efficacy	Safety
<ul style="list-style-type: none"> <li>• Thrombolytic dosing</li> <li>• Thrombolytic infusion duration</li> <li>• Adjuvant therapy</li> <li>• Echocardiogram               <ul style="list-style-type: none"> <li>- Change in RV:LV ratio from baseline</li> <li>- Tricuspid annular plane systolic excursion (TAPSE)</li> <li>- IVC collapse</li> <li>- Estimated right ventricular systolic pressure (RVSP)</li> </ul> </li> <li>• Healthcare utilization: ICU and hospital length of stay (LOS)</li> <li>• Quality of life as measured by PEemb-QoL and EQ-5D-5L VAS – 365 days</li> </ul>	<ul style="list-style-type: none"> <li>• Recurrent VTE</li> <li>• Major bleeding</li> <li>• Mortality</li> <li>• Diagnosis of chronic thromboembolic pulmonary hypertension (CTEPH)</li> </ul>

KEY RESULTS SAFETY (within 30 days)	0 ICH (0/489)	2.5% ISTH Major Bleeding (12/489)	PE: 0.8% (4/489)	Recurrent VTE Confirmed Post-Procedure DVT: 0.2% (1/489)
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<b>Major Bleeding within 30 days</b>	<b>12 (2.5)</b>	Lower Gastrointestinal Hemorrhage	1 (0.2)
Anemia	1 (0.2)	Retinal Hemorrhage	1 (0.2)
Gastrointestinal Hemorrhage	1 (0.2)	Retroperitoneal Hemorrhage	1 (0.2)
Peritoneal Hemorrhage	1 (0.2)	Subdural Hematoma (pre-existing)	1 (0.2)
Procedural Hemorrhage	1 (0.2)	Vascular Access Site Hematoma	1 (0.2)
Laceration	1 (0.2)	<b>Recurrent VTE within 30 days</b>	
Compartment Syndrome	1 (0.2)	Pulmonary Embolism	4 (0.8)
Hematoma	1 (0.2)	Confirmed Post-Procedure DVT	1 (0.2)*

\*34 DVTs were reported, however; 33 of the 34 reported DVTs were identified on duplex ultrasound within 1-2 days of the USCDT procedure and were not differentiated from preexisting DVTs. 30 of the 34 DVTs were reported from one center.

# KNOCOUT PE

## KEY RESULTS | PROCEDURAL

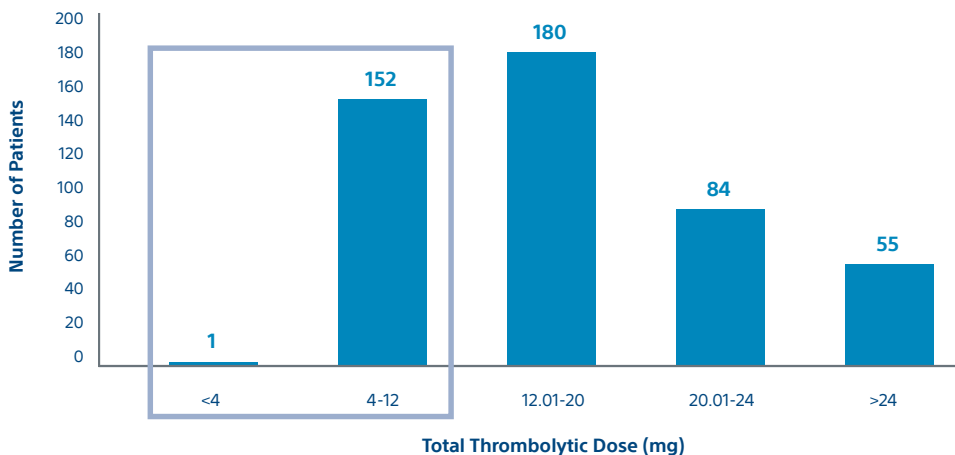
### PROCEDURAL

- Mean dose of r-tPA: 17.9mg (SD 7.3)
- Mean infusion time: 10.4 hrs (SD 5.2)
- Mean time in ICU = 48.9hrs (SD 47.4)

32.4% of patients received < 12mg r-tPA

70.6% of patients received < 20mg r-tPA

### TOTAL THROMBOLYTIC DOSE

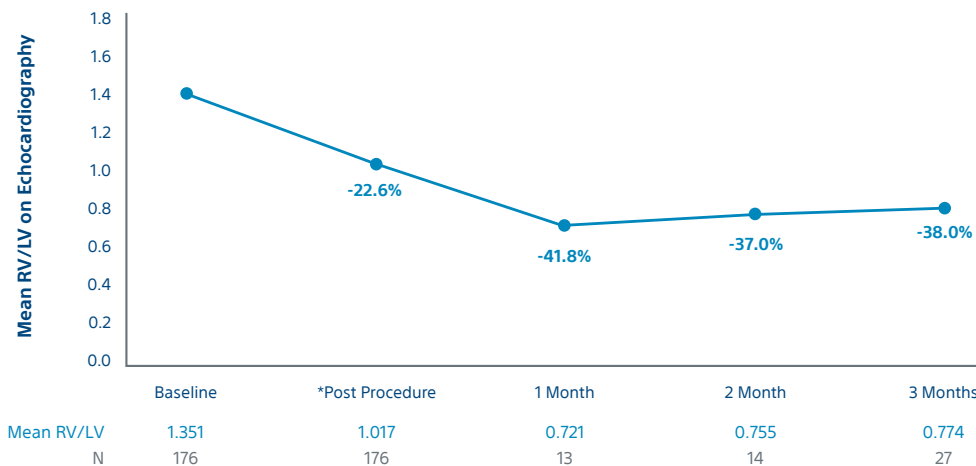


## KEY RESULTS | EFFICACY

### EFFICACY

- RV/LV Reduction from baseline
  - Post-Procedure: 23%
  - 3-months: 38%

### CHANGE IN RV/LV



Significant decrease post-procedure that was sustained over time (P<0.0001)

## KEY RESULTS | QUALITY OF LIFE

### QUALITY OF LIFE

- PEEmb-QOL reduction at 3-months: 41%

QoL Measure	Post-Procedure Mean (SD)	3-Months Mean (SD)	Percent Change Mean (SD)	2-sided P-value
PEEmb-QOL	38.5 (2.1)	16.0 (17.7)	41.1 (114.1)	<0.0001
VAS	63.1 (23.0)	75.5 (19.8)	56 (255.0)	0.0007

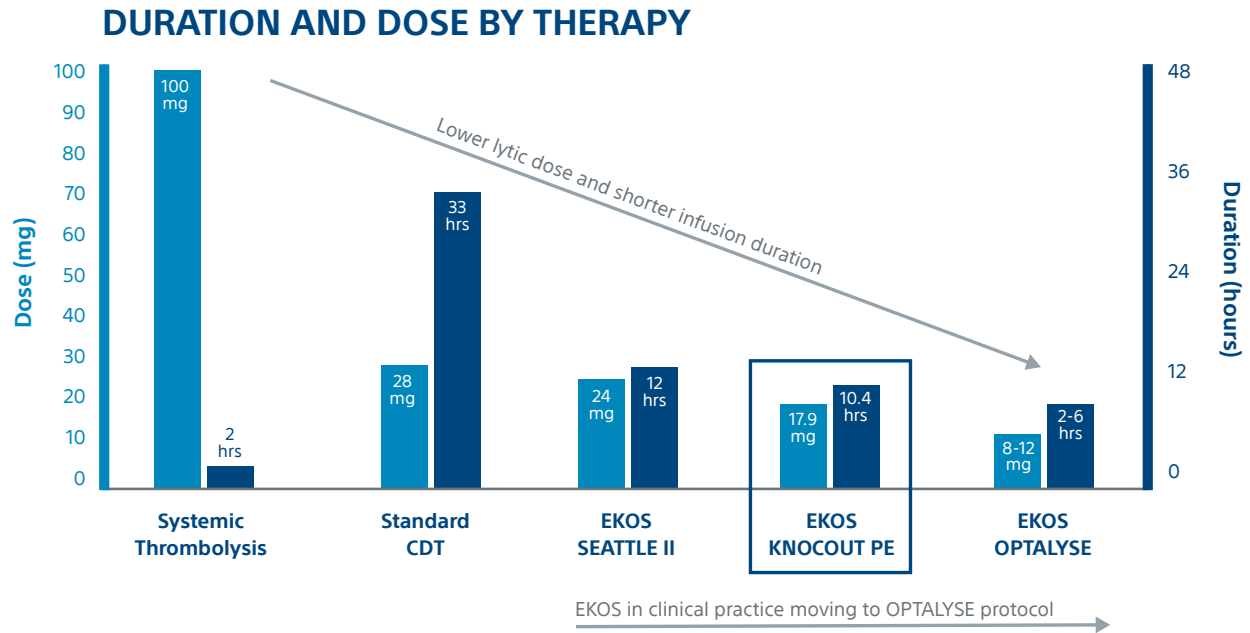
# KNOCOUT PE

## KNOCOUT PE CONCLUSIONS:

Results from this prospective multicenter registry reflect contemporary practice and demonstrate the performance of EKOS in the management of PE with lower total r-TPA dose and shorter infusion duration, marked clinical improvement in RV/LV ratio and low rates of major hemorrhagic complications with no intracerebral hemorrhagic events.

## THROMBOLYTIC THERAPIES:

KNOCOUT PE shows that contemporary clinical practices are moving to low-dose, short duration OPTALYSE protocols. It adds to the growing evidence that EKOS is effective at treating intermediate risk and high risk PE with lower lytic doses and shorter infusion durations compared to other thrombolytic therapies.



**Systemic Thrombolysis** – Konstantinides S, Geibel A, Heusel G, et al. Heparin plus alteplase compared with heparin alone in patients with submassive pulmonary embolism. *N Engl J Med.* 2002;347:1143–1150.

**Standard CDT** – Kuo W et al. *CHEST* 2015; 148(3):667-673.

**SEATTLE II** – Piazza G et al. A Prospective, Single-Arm, Multicenter Trial of Ultrasound-Facilitated, Catheter-Directed, Low-Dose Fibrinolysis for Acute Massive and Submassive Pulmonary Embolism. The SEATTLE II Study. *J Amer Coll Cardiol: Cardiovasc Interventions* 2015; 8(10):1382-1392.

**KNOCOUT** – 3-Month Prospective KNOCOUT Data Presentation at VIVA 2021.

**OPTALYSE** – Tapson VF, Sterling K, Jones N, et al. A randomized trial of the optimum duration of acoustic pulse thrombolysis procedure in acute intermediate-risk pulmonary embolism: the OPTALYSE PE trial. *JACC Cardiovasc Interv.* 2018;11:1401-1410. doi: 10.1016/j.jcin.2018.04.008

#### EKOS Acoustic Pulse Thrombolysis Treatment

**CAUTION:** The law restricts these devices to sale by or on the order of a physician. Indications, contraindications, warnings and instructions for use can be found in the product labelling supplied with each device. Products shown for INFORMATION purposes only and may not be approved or for sale in certain countries. This material is not intended for use in France. Results from different clinical investigations are not directly comparable. Information provided for educational purposes only. PI-1140510-AA

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