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): January 11, 2021

AngioDynamics, Inc.

(Exact Name of Registrant as Specified in Charter)

Delaware	000-50761	11-3146460	
(State or Other Jurisdiction of Incorporation)	(Commission File Number)	(IRS Employer Identification No.)	
14 Plaza D	Prive Latham,New York	12110	
(Address of F	Principal Executive Offices)	(Zip Code)	
	(518) 795-1400		
(Registra	nt's telephone number, including a	rea code)	
Check the appropriate box below if the Form registrant under any of the following provision		usly satisfy the filing obligation of the	
$\hfill\Box$ Written communications pursuant to Rule	425 under the Securities Act (17 C	FR 230.425)	
□ Soliciting material pursuant to Rule 14a-12	2 under the Exchange Act (17 CFR	240.14a-12)	
□ Pre-commencement communications purs	suant to Rule 14d-2(b) under the E	xchange Act (17 CFR 240.14d-2 (b))	
□ Pre-commencement communications purs	suant to Rule 13e-4(c) under the E	xchange Act (17 CFR 240.13e-4 (c))	
Securities registered pursuant to Section 2	L2(b) of the Act:		
Title of each class	<u>Trading Symbol(s)</u>	Name of each exchange on which registered	
Common Stock, par value \$0.01 per share	ANGO	NASDAQ Global Select Market	
Indicate by check mark whether the registrant 1933 (§230.405 of this chapter) or Rule 12b-2			Act of
Emerging growth company \square			
If an emerging growth company, indicate by c for complying with any new or revised financial Act. \Box			

Item 7.01 - Regulation FD Disclosure.

On January 11, 2021, James C. Clemmer, President and Chief Executive Officer of AngioDynamics, Inc. ("AngioDynamics"), will present at the 23rd Annual Needham Virtual Growth Conference. The presentation slides are furnished herewith as Exhibit 99.1.

The presentation slides furnished pursuant to Item 7.01 of this Form 8-K (including Exhibit 99.1) 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 under that Section. Furthermore, the presentation slides shall not be deemed to be incorporated by reference into any filing under the Securities Act of 1933, as amended, or the Exchange Act.

Forward-Looking Statements

This document and its attachments contain forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995. All statements regarding AngioDynamics' expected future financial position, results of operations, cash flows, business strategy, budgets, projected costs, capital expenditures, products, competitive positions, growth opportunities, plans and objectives of management for future operations, as well as statements that include the words such as "expects," "reaffirms," "intends," "anticipates," "plans," "believes," "seeks," "estimates," "optimistic," or variations of such words and similar expressions, are forward-looking statements. These forward-looking statements are not guarantees of future performance and are subject to risks and uncertainties. Investors are cautioned that actual events or results may differ from AngioDynamics' expectations. Factors that may affect the actual results achieved by AngioDynamics include, without limitation, the scale and scope of the COVID-19 global pandemic, the ability of AngioDynamics to develop its existing and new products, technological advances and patents attained by competitors, infringement of AngioDynamics' technology or assertions that AngioDynamics' technology infringes the technology of third parties, the ability of AngioDynamics to effectively compete against competitors that have substantially greater resources, future actions by the FDA or other regulatory agencies, domestic and foreign health care reforms and government regulations, results of pending or future clinical trials, overall economic conditions, the results of on-going litigation, challenges with respect to third-party distributors or joint venture partners or collaborators, the results of sales efforts, the effects of product recalls and product liability claims, changes in key personnel, the ability of AngioDynamics to execute on strategic initiatives, the effects of economic, credit and capital market conditions, general market conditions, market acceptance, foreign currency exchange rate fluctuations, the effects on pricing from group purchasing organizations and competition, the ability of AngioDynamics to integrate purchased businesses, as well as the risk factors listed from time to time in AngioDynamics' SEC filings, including but not limited to its Annual Report on Form 10-K for the year ended May 31, 2020 and its Quarterly Report on Form 10-Q for the fiscal period ended August 31, 2020. AngioDynamics does not assume any obligation to publicly update or revise any forward-looking statements for any reason.

Item 9.01 - Financial Statements and Exhibits.

(d) Exhibits.

Exhibit No. Description

99.1 Presentation slides for the 23rd Annual Needham Virtual Growth Conference, dated January 11, 2021

SIGNATURE

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.

ANGIODYNAMICS, INC. (Registrant)

Date: January 11, 2021 By: /s/ Stephen A. Trowbridge

Name: Stephen A. Trowbridge
Title: Executive Vice President

and

Chief Financial Officer



Forward-Looking Statement

Notice Regarding Forward-Looking Statements

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In the United States, the NanoKnife System has received a 510(k) clearance by the Food and Drug Administration for use in the surgical ablation of soft tissue, and is similarly approved for commercialization in Canada, the European Union and Australia. The NanoKnife System has not been cleared for the treatment or therapy of a specific disease or condition.

Notice Regarding Non-GAAP Financial Measures

Management uses non-GAAP measures to establish operational goals and believes that non-GAAP measures may assist investors in analyzing the underlying trends in AngloDynamics' business over time. Investors should consider these non-GAAP measures in addition to, not as a substitute for or as superior to, financial reporting measures prepared in accordance with GAAP. In this presentation, AngloDynamics has reported adjusted let income before interest, taxes, depreciation and amortization and stock-based compensation); adjusted ent income; adjusted earnings per share and free cash flow. Management uses these measures in its internal analysis and review of operational performance. Management believes that these measures provide investors with useful information in comparing AngloDynamics' performance over different periods. By using these non-GAAP measures, management believes that investors are the picture of the performance of AngloDynamics' underlying business. Management encourages investors to review AngloDynamics' financial results prepared in accordance with GAAP to understand AngloDynamics' performance taking into account all relevant factors, including those that may only occur from time to time but have a material impact on AngloDynamics' financial results. Please see the tables that follow for a reconciliation of non-GAAP measures to measures prepared in accordance with GAAP.



ANGIODYNAMICS

Strategic Transformation



PURSUIT OF LARGER MARKETS

Active portfolio management enables us to compete in larger, growing markets relying on technology & innovation to produce measurable patient outcomes

FOCUSED RESOURCE DEVELOPMENT

Resource deployment is focused in areas that offer best opportunities for success

PORTFOLIO TRANSFORMATION

Portfolio transformation & strength is driven by R&D, M&A, and Clinical & Regulatory

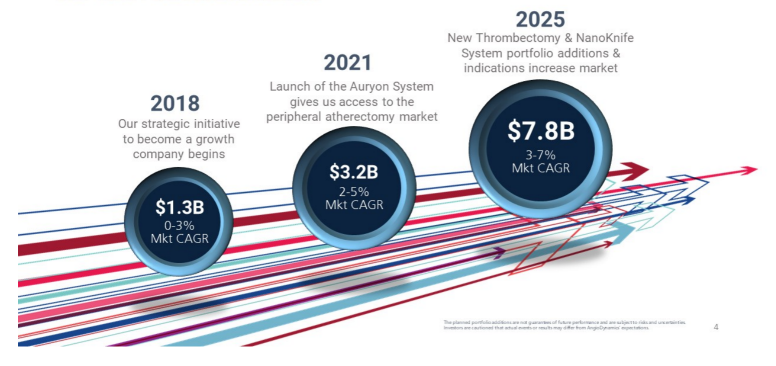
TOP TALENT

Portfolio combined with talent drives value



FOCUSED TRANSFORMATION

U.S. Total Addressable Markets



INVEST FOR GROWTH

Peripheral Atherectomy

AURYON

Thrombus Management

AngioVac Uni-Fuse+

Irreversible Electroporation

NancKnife 3.0





MAINTAIN POSITIONING

Vascular Access Catheters and Accessories

Diagnostic Catheters, Guidewires and Kits

Endovenous Laser Treatment

Microwave & Radiofrequency Tumor Ablation

Lung Biopsy Safety

Radiation Treatment Stabilization Balloons

FOCUSED INVESTMENT











Leveraging 3 main drivers to carve out our space in large, growing markets through innovative, disruptive technologies that treat patients with cancer, promote healthy blood flow and deliver critical

M & A

therapies.

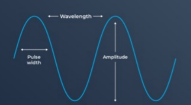
R&D

Clinical and Regulatory Pathway Expansion

















Z.Omm

Aspiration capability and indicated
Aspiration and Off-Center capabilities^{for} Peripheral Atherectomy and ISR and indicated for Peripheral
Atherectomy and
In-Stent Restenosis (ISR)

Why wavelength matters

Each type of tissue interacts differently with a given wavelength

produces a photon energy but high enough to vaporize calcium.d, g

Why pulse width and amplitude matter

Greater amplitude is achieved with shorter pulses, which can deposit energy before thermal diffusion occurs

the lesion and spare the vessel.c

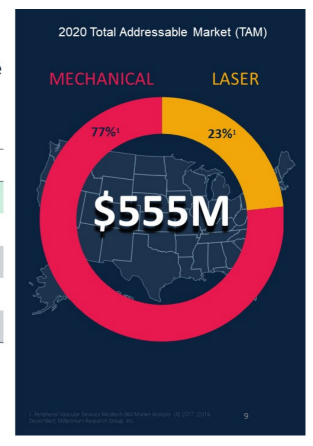


c, d, g See appendix for references

PERIPHERAL ATHERECTOMY

US Addressable Markets & Competitive Landscape

Company	Product	Thrombus	Plaque	Fibrotic	ISR	Calcified
ANGO	Auryon	***	***	**	***	**
Philips	Excimer Laser	**	**		***	*
MDT	TurboHawk		***	*		*
BSC	Jetstream	**	*	*		
CSI	Diamonback 360		**	*		***





PERIPHERAL ATHERECTOMY

Continuing our momentum of growth

CLINICAL COMPENDIUM

- Expand awareness by creating a strong podium and clinical publication presence
- Clinically differentiated validation through new users, data collection (PATHFINDER I & II), and clinical outcomes







- Targeted commercial & clinical expansion
- · Meaningful physician training programs
- Increase physician ambassador programs



PRODUCT DEVELOPMENT

- · Product development pipeline
- Portfolio and product indication expansion
- · Continuous voice of customer



Deep Vein **Thrombosis**



Pulmonary Embolism



A blood clot that forms in a deep vein, usually the leg, groin or arm

A DVT breaks free from a vein wall and travels to the lungs blocking some or all of the blood supply

208,000 Iliofemoral Cases1

171,000 High-risk & intermediate-risk PE Cases1





Clot in Transit (traveling through the heart)

Clot in Pulmonary Arteries (PE)



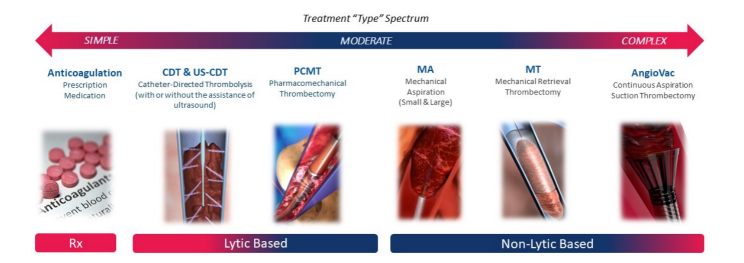
Venous Thromboembolism

DVT and PE are collectively referred to as VTE

100,000 - 300,000 VTE-Related Deaths in the USA Annually²

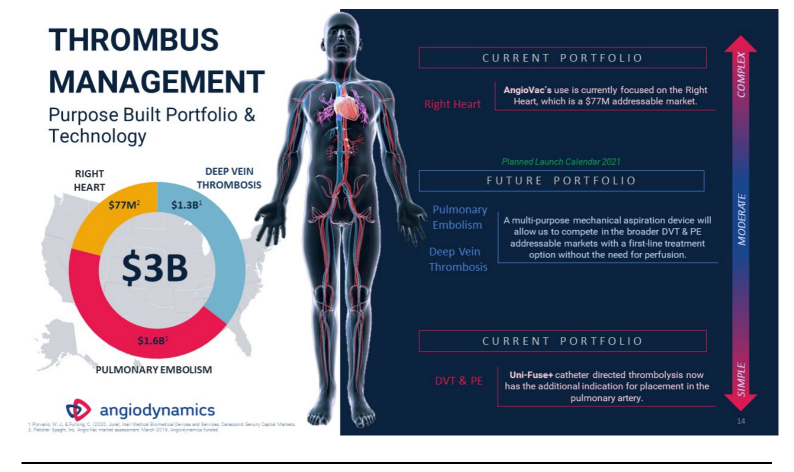
DVT & PE TREATMENT OPTIONS

Percutaneous Thrombectomy





Illustrations and Images not Produced by AngloDynamics Induc https://you.ubu/DQUIDSBRick https://www.perumbninc.com/indgolightning/ https://www.weculamedcure.com/pugeoducis





Difference

The AngioVac System allows for the **continuous aspiration**of embolic material such as fresh, soft thrombi or vegetation from the venous system

Utilizing a self-expanding, nitinol reinforced funnel tip

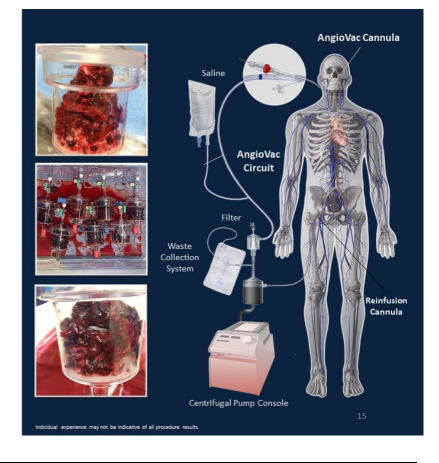
Simultaneously reinfusing the patient's own filtered blood to limit procedural blood loss





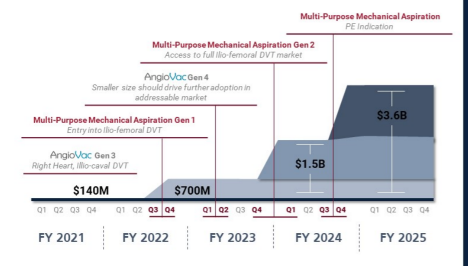






THROMBUS MANAGEMENT

Planned Portfolio Additions & U.S. Addressable Markets Expansion



THE NEXT PORTFOLIO INNOVATION

A purpose-built, innovative product leveraging the strengths of the AngioVac cannula technology with off-circuit manual aspiration control



Powerful

Proven, funnel tip design allows efficient aspiration and compression of large clot burden



Controlled

Designed to allow the end-user command and control of the

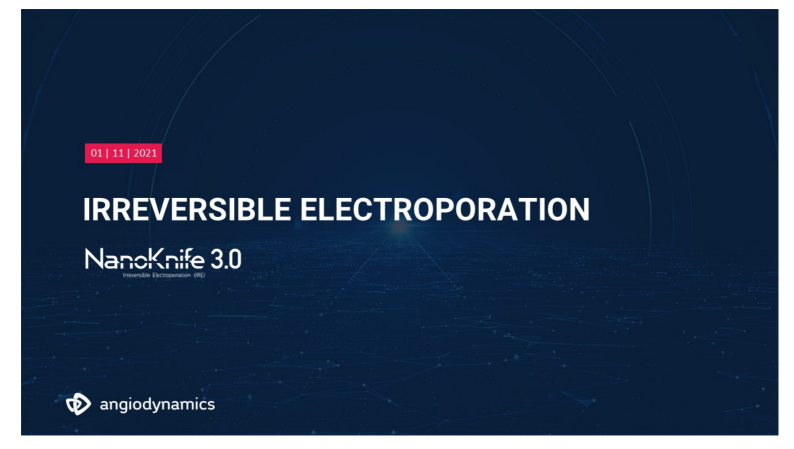


Versatile

Broadens our Thrombus Management portfolio and designed to provide an intuitive, first-line treatment option without the need for lytics and advanced procedural support

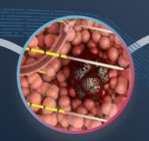
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The planned portfolio additions are not guarantees of future performance and are subject to risks and uncertainties. Investors are cautioned that actual events or results may differ from Angio Dynamics' expectations.



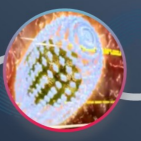
IRREVERSIBLE ELECTROPORATION

NanoKnife



Probe placement

Placement of probes around a target area provides effective treatment coverage a-d



Electrical field

Electrical pulses cause pores to form within the cellular membrane leading to nonthermal cell death ^{a-d}



Decellularization

Target tissue undergoes complete decellularization while preserving the underlying structure of blood vessels, nerves, ducts, and tissue ad



Non-thermal

By preserving those underlying structures the potential for revascularization of treated tissue is maintained ^{e-d}



Revascularization

The NanoKnife System's unique technology enables physicians to provide localized treatments in locations and at times previously unavailable to them a-d

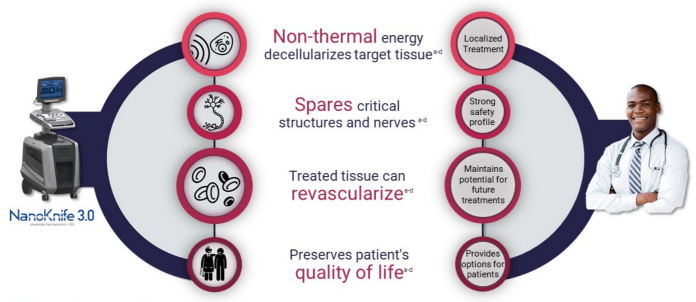


Maor E. et al., The effect of imperable electroparation on blood lessals. Technol. Carner Res. Treat. 6(4), 307–312 (2007), 10.11.177/15330349/07009000407.

Ribinolsy, S. O. Ribino, S. C. et al., and Mikas, P. Herrers ble electroparation a new abstrom modality-crinical implications. Technol. Carner Res. Treat. 6(4), 295–300 (2007), 10.1177/153303400700600106.

Onk G. Wikas P., and Rubinsky B., kneversible electroparation; implications for prostate abstration. Technol. Carner Res. Treat. 6(4), 295–300 (2007), 10.1177/153303400700600405.

NANOKNIFE PROVIDES INNOVATION DOCTORS NEED



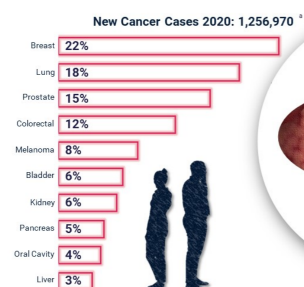


aor E. et al., The effect of inversible electroporation on blood wassels, Technol, Cancer Res. Treats. 6(4), 307–312 (2007).10.1177/15.8303.460700.600407).

doinisty, B., Onlik, G., and Mikus, P., Inversible electroporation: a new ablation modality—clinical implications. Technol. Cancer Res. Treat. 6, 47–49. (2007). doi:10.1177/15.3303.4607006.0010.6.

ink G., Mikus P., and Rubinsky B., Inversible electroporation: implications for prostate ablation, Technol. Cancer Res. Treat. 6(4), 297–300 (2007).10.1177/15.3303.4607006.0010.6.









This safety study will evaluate the use of the NanoKnife System as a focal therapy option for prostate tissue.



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CLINICAL EXPANSION

Within the U.S. 191,930 men will be diagnosed with prostate cancer in 2020. 27% will undergo radiotherapy of which 66% report E.D. at 36 months

31% will undergo a radical prostatectomy of which **79%** report E.D. at 36 months b.c

42% will undergo active surveillance

An estimated **38%** of these patients are ideal candidates for focal therapy_d



Focal therapy

bridges the gap between whole gland treatment and active surveillance in an attempt to maximize quality of life

by avoiding the effects of whole gland radiation or surgery.

However...
adoption remains low
(<5%) due to gaps
within existing technology

within existing technology.





NANOKNIFE PLATFORM



Expanding Indications Pancreas pivotal study underway (DIRECT) Prostate safety study underway FDA Pre-Sub meeting regarding prostate tissue indication complete



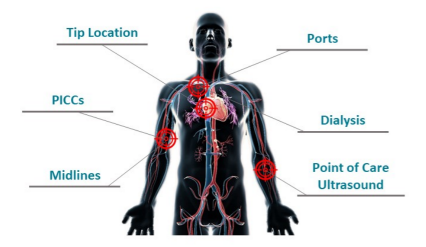






VASCULAR ACCESS

Safely delivering medication to patients











PORTFOLIO

Delivering on our product road map including development of an integrated navigation & tip location technology that enables the use of our BioFlo PICC.

MARKET ACCESS

Maximize clinical differentiation by reducing thrombus accumulation through the utilization of our BioFlo family of



PERFORMANCE

Maintain a strong culture of execution and collaboration through disciplined sales & marketing plans

First Half FY2021 Highlights

Financial Performance

\$ in Millions (except per share data)	1H FY2021	1H FY2020	YOY Change
Revenue	\$143.0M	\$136.0M	5.1%
Gross Margin	53.1%	58.6%	(550 bps)
Adjusted EPS	\$0.03	\$0.14	(\$0.11)
Adjusted EBITDA	\$9.6	\$13.7	\$4.1
Free Cash Flow	\$2.8	(\$4.6)	\$7.4

\$ Millions	Q2 FY2021	Q4 FY2020	YOY Change
Cash	\$58.0	\$54.4	\$3.6
Debt	\$40.0	\$40.0	\$-



FY 2021 Guidance (Unchanged)			
Revenue	\$278 - \$284		
Adjusted EPS	\$0.00 - \$0.05		

Growth From Key Products

Key Product Categories	Q1 FY2021	Q2 FY2021	1H FY2021
AngioVac®	46%	24%	34%
NanoKnife® Disposables	(5%)	30%	12%
Auryon*	\$1.1	\$2.1	\$3.2

^{*} Auryon reflects revenue contribution in quarter vs growth rate due to acquisition date in FY20 $\,$

ANGIODYNAMICS

Strategic Transformation



PURSUIT OF LARGER MARKETS

Active portfolio management enables us to compete in larger, growing markets relying on technology & innovation to produce measurable patient outcomes

FOCUSED RESOURCE DEVELOPMENT

Resource deployment is focused in areas that offer best opportunities for success

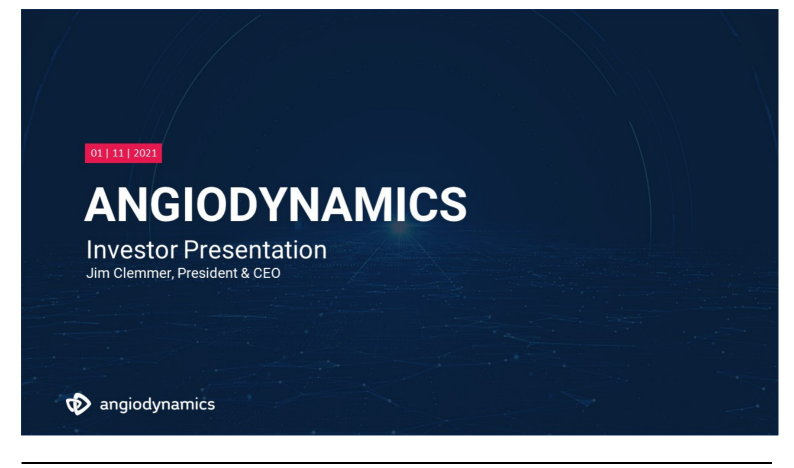
PORTFOLIO TRANSFORMATION

Portfolio transformation & strength is driven by R&D, M&A, and Clinical & Regulatory

TOP TALENT

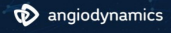
Portfolio combined with talent drives value





• **Peripheral Atherectomy References:

a. Rundback J, Chandra P, Brodmann M, Weinstock B, Sedillo G, Cawich I, et al. Novel laser-based catheter for peripheral atherectomy: 6-month results from the Eximo Medical B-Laser™ IDE study. Catheter Cardiovasc Interv. 2019;1-8. b. Shammas NW, Chandra P, Brodmann M, Weinstock B, Sedillo G, Cawich I, et al. Acute and 30-day safety and effectiveness evaluation of Eximo Medical's B-Laser™, a novel atherectomy device, in subjects affected with infrainguinal peripheral arterial disease: Results of the EX-PAD-03 trial. Cardiovas Revasc Med. 2020;21(1):86-92. c. Auryon. Instructions for use. AngioDynamics; 2019. d. Herzog A, Bogdan S, Glikson M, Ishaaya AA, Love C. Selective tissue ablation using laser radiation at 355 nm in lead extraction by a hybrid catheter, a preliminary report. Lasers Surg Med. 2016;48(3):281-287. c. Herzog A, Steinberg I, Gaisenberg E, Nomberg R, Ishaaya AA. A route to laser angioplasty in the presence of fluoroscopy contrast media, using a nanosecond-pulsed 355-nm laser. IEEE J Sel Top Quantum Electron. 2016;22(3):342-347. f. Kuczmik W, Kruszyna L, Stanisic MG, Dzieciuchowicz L, Ziaja K, Zelawski W, et al. Laser atherectomy using the novel B-Laser™ catheter, for the treatment of femoropopliteal lesions: twelve-month results from the EX-PAD-01 study. Not yet published. g. Spectranetics Corporation. CVX-300 Excimer Laser System: Operator's Manual. Version 28. 2019:1-56.



AngioVac Cannula Indication for Use: The AngioVac Cannula is indicated for use as a venous drainage cannula and for removal of fresh, soft thrombi or emboli during extracorporeal bypass for up to 6 hours. | Contraindications: Contraindicated for patients with severe arterial or venous vascular disease, contraindicated for removal of chronic firmly adherent intravascular material (e.g., atherosclerotic plaque, chronic pulmonary embolism) and for use in the right heart or pulmonary arteries during active cardiopulmonary resuscitation. | Refer to Directions for Use and/or User Manual provided with the product for complete Instructions, Warnings, Precautions, Possible Adverse Effects and Contraindications prior to use of the product. | AngioVac Circuit Indications for Use + AngioVac Circuit is indicated for use in product. | AngioVac Circuit Indications for Use and/or User Manual provided with the product for complete Instructions, Warnings, Precautions, Possible Adverse Effects and Contraindications. Refer to Directions for Use and/or User Manual provided with the product for complete Instructions, Warnings, Precautions, Possible Adverse Effects and Contraindications prior to use of the product. | CAUTION: Federal law (USA) restricts this device to sale by or on the order of a physician. | Refer to Directions for Use and/or/User Manual provided with the product for complete Instructions, Warnings, Precautions, Possible Adverse Effects and Contraindications prior to use of the product. | NanoKnife System Indications For Use and Important Risk Information US: The NanoKnife System with six outputs is indicated for surgical ablation of soft tissue. | CE: The NanoKnife System Indications For Use and Important Risk Information US: The NanoKnife System with six outputs is indicated for surgical ablation of soft tissue. | CE: The NanoKnife System is a medical device for cell membrane electroporation is a phenomenon that occurs in cell membranes as cells are exposed to an electrical field of sufficiently high intensity. Th

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