

PERRY JOHNSON LABORATORY ACCREDITATION, INC.

Certificate of Accreditation

Perry Johnson Laboratory Accreditation, Inc. has assessed the Laboratory of:

Biomedical Device Consultants & Laboratories, LLC (BDC Laboratories) 4060 Youngfield St., Wheat Ridge, CO 80033

(Hereinafter called the Organization) and hereby declares that Organization is accredited in accordance with the recognized International Standard:

ISO/IEC 17025:2017

This accreditation demonstrates technical competence for a defined scope and the operation of a laboratory quality management system (as outlined by the joint ISO-ILAC-IAF Communiqué dated April 2017):

Electrical, Dimensional Inspection and Mechanical Testing (As detailed in the supplement)

Accreditation claims for such testing and/or calibration services shall only be made from addresses referenced within this certificate. This Accreditation is granted subject to the system rules governing the Accreditation referred to above, and the Organization hereby covenants with the Accreditation body's duty to observe and comply with the said rules.

For PJLA:

Diacy Szenspen

Tracy Szerszen President

Perry Johnson Laboratory Accreditation, Inc. (PJLA) 755 W. Big Beaver, Suite 1325 Troy, Michigan 48084

Initial Accreditation Date:	Issue Date:	Expiration Date:
September 29, 2015	December 4, 2021	December 31, 2023
Revision Date:	Accreditation No: 87132	<i>Certificate No:</i>
October 14, 2022	87132	L21-741-R1

The validity of this certificate is maintained through ongoing assessments based on a continuous accreditation cycle. The validity of this certificate should be confirmed through the PJLA website: <u>www.pjlabs.com</u>



Certificate of Accreditation: Supplement

Biomedical Device Consultants & Laboratories, LLC (BDC Laboratories)

4060 Youngfield St., Wheat Ridge, CO 80033 Contact Name: Devin McBlair Phone: 303-456-4665

Accreditation is granted to the facility to perform the following testing:

FIELD OF TEST	ITEMS, MATERIALS OR PRODUCTS TESTED	SPECIFIC TESTS OR PROPERTIES MEASURED	SPECIFICATION, STANDARD METHOD OR TECHNIQUE USED	RANGE (WHERE APPROPRIATE) AND DETECTION LIMIT
Electrical ^F	Implantable Medical Devices	Electrochemical Corrosion, Cyclic Potentiodynamic	ASTM F2129 Lab Developed Method TM- 0080	± 600 mA, ± 800 mV
		Electrochemical Corrosion, Galvanic	ASTM F3044 Lab Developed Method TM- 0083	± 600 mA, ± 800 mV
Mechanical ^F	Endovascular Devices, Vascular Prostheses	Pulsatile Durability	ASTM F2477 ISO 25539-1, Annex D.5.2.3.2 ISO 25539-2, Annex D.5.3.3.2 Lab Developed Method TM- 0002	0.5 mm to 65 mm
		Determination of Dynamic Compliance	ISO 7198, Annex A.5.9 Lab Developed Method TM- 0001	0.5 mm to 49.5 mm
	Cardiac Valve Prostheses	Hydrodynamic Valve Performance Steady Flow	ISO 5840-1, Annex I Lab Developed Method TM- 0077	Forward Flow Rate: 0 L/min to 35 L/min Back Pressure: 10 mmHg up to 300 mmHg
		Hydrodynamic Valve Performance Pulsatile Flow	ISO 5840-2, Annex F ISO 5840-3, Annex C Lab Developed Method TM- 0013	Flow Rate: 0 L/min to 10 L/min Driving Waveform: Sinusoidal, Arbitrary Frequency: 2 bpm to 240 bpm
		Durability Testing	ISO 5840-1, Annex J Lab Developed Method TM- 0014	Excitation Waveforms: Sine & Modified Sine, Arbitrary Operating Frequency: 5 Hz to 50 Hz
	Medical Devices	Acute Particulate Matter Generation and Coating Integrity	ASTM F2743 ASTM F3320 Lab Developed Methods TM- 0022 and TM-0081	2 μm to 1 000 μm
		Examination Testing: Guidewire Flex, Guidewire Fracture, Coating Integrity	ISO 25539-3 Annex D.5.2.8 ISO 10555-1 Clause 4.4 ISO 10555-3 Clause 4.2 ISO 10555-6 Clause 4.2. ISO 11070 Clause 4.3 ISO 11070 Annex F ASTM F2743 ASTM F3320	Up to 100 X
			Lab Developed Method TM- 0035	



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Mechanical ^F	Medical Devices	Dimensional Verification	ISO 25539-1 Annex D.5.1.2 ISO 25539-1 Annex D.5.2.7.1 ISO 25539-2 Annex D.5.2.3 ISO 25539-2 Annex D.5.3.5.1 ISO 25539-3 Annex D.5.1.1 ISO 25539-3 Annex D.5.5.1 ISO 25539-3 Annex D.5.2.3 ISO 10555-1 Clause 5 Lab Developed Method TM-0041	Up to 152 mm
		Visual Inspections	ISO 25539-3 Annex D.5.2.8 ISO 10555-1 Clause 4.4 ISO 10555-3 Clause 4.2 ISO 10555-6 Clause 4.5.2 ISO 11070 Clause 4.3 Lab Developed Method TM-0035	Up to 100 X
	Endovascular Prostheses, Vascular Stents	Kink Radius	ISO 25539-1 Annex D.5.2.5.5 ISO 25539-2 Annex D.5.3.4.5 ISO 7198 Annex A.5.8 ASTM F3505 Lab Developed Method TM-0030	Up to 200 mm
	Endovascular Prostheses, Vascular Stents	Torsional Durability	ISO 25539-1 Annex D.5.2.3.6 ISO 25539-2 Annex D.5.3.3.5 ASTM F2942 Lab Developed Method BDC-TM- 0082	Angle of twist -10 to 90°
	Guidewires	Guidewire Flex	ISO 11070 Annex G Lab Developed Method TM-0035	Up to 100 X
	Guidewires Coated Medical Devices	Guidewire Fracture Coating Integrity Inspection	ISO 11070 Annex F Lab Developed Method TM-0035 ASTM F2743 ASTM F3320 Lab Developed Method TM-0035	-
	Implantable Medical Devices ^{FO}	Measurement of Radio Frequency Induced Heating	ASTM F2182 Lab Developed Method TM-0053	Field Strengths: 1.5 T, 3 T
		Artifacts from Passive Implants	ASTM F2119 Lab Developed Method TM-0055	-
		Magnetically Induced Torque Measurement of	Lab Developed Method TM-0052	-
		Magnetically Induced Displacement Force	Lab Developed Method TM-0054	

This supplement is in conjunction with certificate #L21-741-R1



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Dimensional	Endovascular	Length to Diameter	ISO 25539-1 Annex D.5.2.7.3	Up to 152 mm
Inspection ^F	prostheses,	Relationship	ISO 25539-2 Annex D.5.3.5.3	1
-	Vascular stents	-	Lab Developed Method TM-0041	
		Elastic Recoil	ISO 25539-1 Annex D.5.2.7.4]
			ISO 25539-2 Annex D.5.3.5.4	
			ASTM F2079	
			Lab Developed Method TM-0041	
	Endovascular	Diameter to Balloon	ISO 25539-1 Annex D.5.2.7.2	
	prostheses,	Inflation Pressure	ISO 25539-2 Annex D.5.3.5.2	
	Vascular stents,		ISO 10555-4 Annex D	
	Balloon dilatation		ASTM F2081	
	catheters		Lab Developed Method TM-0041	
	Medical Devices	Dimensional	ISO 25539-1 Annex D.5.1.2	
		verification	ISO 25539-1 Annex D.5.2.7.1	
			ISO 25539-2 Annex D.5.2.3	
			ISO 25539-2 Annex D.5.3.5.1	
			ISO 25539-3 Annex D.5.1.1	
			ISO 25539-3 Annex D.5.5.1	
			ISO 25539-3 Annex D.5.2.3	
			ISO 10555-1 Clause 5	
			Lab Developed Method TM-0041	
			ASTM F2081	
	Vascular stents	Profile Effect/Flaring	ISO 25539-2 Annex D.5.2.7	
			Lab Developed Method TM-0041	
			ASTM F2081	
		Dogboning	ISO 25539-2 Annex D.5.2.2.5	
			Lab Developed Method TM-0041	
	Vascular prostheses	Microscopic Porosity	ISO 7198, Annex A.5.1.1.3	
		(inter-nodal distance)	Lab Developed Method TM-0041	
		Usable Length/Width	ISO 7198 Annex A.5.3	
			Lab Developed Method TM-0041	
		Relaxed Inner	ISO 7198 Annex A.5.4	
		Diameter	Lab Developed Method TM-0041	
		Wall Thickness	ISO 7198 Annex A.5.6	Up to 13 mm
			Lab Developed Method TM-0041	

- 1. The presence of a superscript F means that the laboratory performs testing of the indicated parameter at its fixed location. Example: Mechanical^F would mean that the laboratory performs this testing at its fixed location.
- 2. The presence of a superscript FO means that the laboratory performs testing of the indicated parameter both at its fixed location and onsite at customer locations. Example: Outside Micrometer^{FO} would mean that the laboratory performs this testing at its fixed location and onsite at customer locations.

This supplement is in conjunction with certificate #L21-741-R1