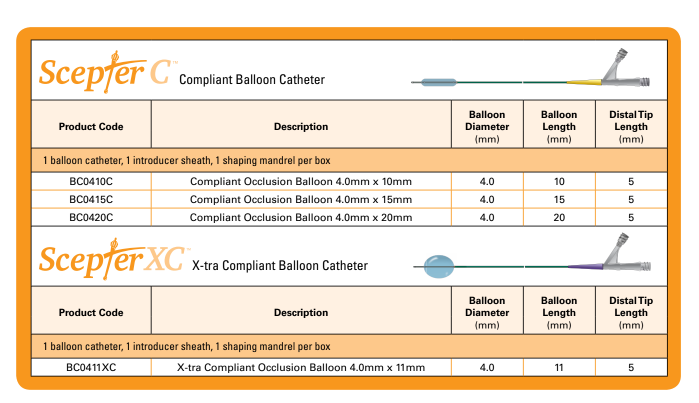




The New Standard of Excellence, Redefining Deliverability, Versatility, and Control



Features Benefits

14 Wire Compatible Provides more choices of guidewire (Traxcess® docking wire compatible)

Improves trackability and stability with better performance of the wire

Hydrophilic Coating on Significantly improves trackability

the Balloon

Enables use of longer balloons without compromising trackability

Soft & Long Tip Improves balloon stability and trackability

Low Profile Ensures compatibility with double catheter technique in 6F guide

Improves trackability

Distal Tip Marker Positive identification of distal tip location



MicroVention, Inc.
Worldwide Headquarters
1311 Valencia Avenue
Tustin, CA 92780 USA
MicroVention UK Limited
MicroVention Europe, S.A.R.L.
MicroVention Deutschland GmbH

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EC-CERTIFICATE



(Full quality assurance system)

This is to certify that the company

MicroVention, Inc.

1311 Valencia Ave. Tustin, CA, 92780 United States of America

has implemented and maintains a full quality assurance system which applies to the products at every stage from design to final controls.

Through an audit, documented in a report, performed by DQS Medizinprodukte GmbH, it was verified that the management system fulfills the requirements of

Annex II – excluding Section 4 of Council Directive 93/42/EEC concerning medical devices

with respect to the following medical devices:

Embolization Prostheses and Accessories, Intravascular Access Devices (Occlusion Balloon Catheters, Guiding and Aspiration Catheters, Microcatheters and Guidewires) and Accessories, Stents, Clot and Foreign Body Retrieval Devices, Liquid Embolic System, Embolic Protection System, Aneurysm Embolization Device, Microspheres, Aspiration Devices, and Detachment Controller Units as listed in Annex.

The manufacturer is subject to surveillance according to Annex II, Section 5. The CE marking with the Notified Body Identification Number (0297) may be affixed on the devices listed in the certificate. An EC Design Examination Certificate according to Annex II, Section 4 is required for class III devices covered by this certificate. The certificate is in the case of class I(s) devices (I(s) = class I products placed on the market in sterile conditions) limited to the aspects of manufacture concerned with securing and maintaining sterile conditions. The certificate is in the case of class I(m) devices (I(m) = class I devices with a measuring function) limited to the aspects of manufacture concerned with the conformity of the products with the metrological requirements.

Certificate registration no. 411133 MR2
Certificate unique ID 170776096
Effective date 2021-04-29
Expiry date 2024-05-26
Frankfurt am Main 2021-04-29

DQS Medizinprodukte GmbH

Sigrid Uhlemann Managing Director Dr. Thomas Feldmann Head of Certification Body

August-Schanz-Straße 21, 60433 Frankfurt am Main, Tel. +49 (0) 69 95427-300, medical.devices@dqs-med.de







Certificate registration No.: 411133 MR2

Certificate unique ID: 170776096

Effective date: 2021-04-29

MicroVention, Inc.

1311 Valencia Ave. Tustin, CA, 92780 United States of America

Production Sites:

MicroVention, Inc.
 Enterprise,
 Aliso Viejo, CA 92656
 United States of America

2.MicroVention, Inc.1311 Valencia Ave.Tustin, CA 92780United States of America

3. MicroVention Costa Rica, S.R.L. Zona Franca Coyol Alajuela, Costa Rica







Certificate registration No.: 411133 MR2

Certificate unique ID: 170776096

Effective date: 2021-04-29

MicroVention, Inc.

1311 Valencia Ave. Tustin, CA, 92780 United States of America

	Device Groups:	Device Family:	Devices:	Risk Class	Production Site
	Embolization Prothese	MicroPlex Coil System (MCS) & HydroCoil Embolic System (HES) with V-Trak Delivery System	MicroPlex 10 Platinum Coil System (MCS) Endovascular Embolization Coil - Cosmos10 - HyperSoft 3D - HyperSoft Helical - Helical 10 - VFC - Compass 10 - Complex 10	III	1,2,3
		System.	MicroPlex 18 Platinum Coil System (MCS) Endovascular Embolization Coil - Cosmos 18 - Helical 18 - Compass 18 - Complex 18	III	1,2,3
			HydroCoil 10 Embolic System (HES) Endovascular Embolization Coil - HydroFrame 10 - HydroSoft Helical - HydroSoft 3D - HydroFill	III	1,2,3
			HydroCoil 18 Embolic System (HES) Endovascular Embolization Coil - HydroFrame 18	III	1,2,3
		AZUR®	AZUR® HydroCoil Detachable Embolization Coils 18 & 35	IIb	1,2,3
		Peripheral Coil System	AZUR® HydroCoil Pushable Embolization Coils 18 & 35 AZUR® Framing Detachable Coils 18 & 35	IIb	1,2,3
				IIb	1,2,3
			AZUR® Injectable Coil System 18 & 35	IIb	1,2,3
			AZUR Detachable 18 AZUR PURE Pushable Coil System 18 & 35	IIb IIb	1,2,3 1,2,3
			AZUR CX Detachable 18 & 35 AZUR Vascular Plug	IIb IIb	1,2,3 1,2,3

This annex is only valid in connection with the above-mentioned certificate.

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Certificate registration No.: 411133 MR2

Certificate unique ID: 170776096

Effective date: 2021-04-29

MicroVention, Inc.

1311 Valencia Ave. Tustin, CA, 92780 United States of America

Device Groups:	Device Family:	Devices:	Risk Class	Production Site
Detachment Controller Units		V-Grip® Detachment Controller V-Grip® PLUS Detachment Controller WEB Detachment Controller AZUR® Detachment Controller	lla Ila Ila Ila	1,2 1,2 1,2 1,2
Intravascular Access Devices (Occlusion Balloon Catheters, Guiding and Aspiration Catheters,	5	Traxcess® 14 Guidewire Traxcess® 14 EX Guidewire Traxcess® 14 SELECT Guidewire Traxcess® 7 Mini Guidewire Traxcess® 7 Mini XSoft Guidewire Traxcess® Docking Wire	 a	1,2 1,2 1,2 1,2 1,2 1,2
Microcatheters and Guidewires	;)	Chaperon® Guiding Catheter System	Ш	2
	,	Headway® 17 Advanced Soft	Ш	1,2,3
		Microcatheter Headway [®] 17 Advanced Microcatheter	Ш	1,2,3
		Headway® 21 Microcatheter Headway® 27 Microcatheter Headway Duo Microcatheter Scepter C™ Occlusion Balloon Catheter Scepter XC™ Occlusion Balloon Catheter Scepter Mini™ Occlusion Balloon Catheter SOFIA™ Distal Access Catheter SOFIA™ Select Catheter SOFIA™ PLUS Catheter SOFIA™ Flow PLUS Catheter SOFIA™ Guiding Catheter SOFIA™ Flow Catheter SOFIA® EX Catheter KANSHAS Drug Coated Balloon		1,2,3 1,2,3 1,2,3 1,2,3 1,2,3 1,2,3 1,2,3 1,2,3 1,2,3 1,2,3 1,2,3 1,2,3 1,2,3 1,2,3 1,2,3
		VIA [™] 17 Microcatheter VIA [™] 21 Microcatheter VIA [™] 27 Microcatheter VIA [™] 33 Microcatheter Wedge Microcatheter PG Pro Microcatheter	 a	1,2 1,2 1,2 1,2 1,2,3 1,2,3







Certificate registration No.: 411133 MR2
Certificate unique ID: 170776096
Effective date: 2021-04-29

MicroVention, Inc.

1311 Valencia Ave. Tustin, CA, 92780 United States of America

Device Groups:	Device Family:	Devices:	Risk Class	Production Site
Stents		LVIS™ Intraluminal Support Device	III	1,2,3
Otents		LVIS™ Jr. Intraluminal Support Device	III	1,2,3
		LVIS™ EVO™ Intraluminal Support Device	Ш	1,2,3
		LVIS™ X™ Intraluminal Support Device	III	1,2,3
		LVIS™ Jr. X™ Intraluminal Support Device	III	1,2,3
		LVIS™ EVO™ X™ Intraluminal Support Device	III	1,2,3
		FRED™ Flow Re-Direction Endoluminal Device	III	1,2,3
		FRED Jr.™ Flow Re-Direction Endoluminal Device	Ш	1,2,3
		FRED X™ Flow Re-Direction Endoluminal Devices	Ш	1,2,3
		FRED OMEGA™ Flow Re-Direction Endoluminal Devices	III	1,2,3
		CASPER™ RX Carotid Artery Stent System	III	1,2,3
		Roadsaver Carotid Artery Stent System	Ш	1,2,3
Peripheral Vascular Stent System		RENZAN™ Peripheral Vascular Stent System	IIb	1,2,3
Clot Retriever		ERIC™ Retrieval Device	Ш	1,2,3
Liquid Embolic System		PHIL™ Liquid Embolic System	Ш	1,2
Microspheres		HydroPearl Microspheres	IIb III	1,2 1,2
		LifePearl Microspheres BioPearl® Microspheres	III	1
This annex is only	valid in connec	ction with the above-mentioned certificate.		5/6







Certificate registration No.: 411133 MR2

Certificate unique ID: 170776096

Effective date: 2021-04-29

MicroVention, Inc.

1311 Valencia Ave. Tustin, CA, 92780 United States of America

Device Groups:	Device Family:	Devices:	Risk Class	Production Site
Embolic Protection Device (EPS)		Empro Embolic Protection System Nanoparasol Embolic Protection System	III III	1,3 1,3
Aneurysm Embolization Device		WEB™ Aneurysm Embolization System	III	1,2
Aspiration Kit		Aspiration Tubing Kit Aspiration Syringe Kit	ls Is	1,2 1,2
BOBBY™ Balloon Guide Catheter		BOBBY™ Balloon Guide Catheter	III	1,2







CERTIFICATE



This is to certify that the company

MicroVention, Inc.

35 Enterprise Aliso Viejo, CA, 92656 United States of America

with the organizational units/sites as listed in the annex

has implemented and maintains a Quality Management System.

Scope:

Design, Development, Manufacturing and Distribution of Embolization Prostheses and Accessories, Intravascular Access Devices (Occlusion Balloon Catheters, Guiding and Aspiration Catheters, Microcatheters and Guidewires), Stents, Clot and Foreign Body Retrieval Devices, Liquid Embolic System, Embolic Protection System, Aneurysm Embolization Device, Microspheres, Aspiration Devices and Detachment Controller Units.

Through an audit, documented in a report, performed by DQS Medizinprodukte GmbH, it was verified that the management system fulfills the requirements of the following standard:

DIN EN ISO 13485: 2016 + AC: 2017-07

EN ISO 13485: 2016 + AC: 2016

ISO 13485: 2016

Certificate registration no. 411133 MP2016

Certificate unique ID 170780788

Effective date 2022-07-07

Expiry date 2024-09-26

Frankfurt am Main 2022-07-07







DQS Medizinprodukte GmbH

J. Ml luca

Sigrid Uhlemann Managing Director Dr. Thomas Feldmann Head of Certification Body



Certificate registration No.: 411133 MP2016

Certificate unique ID: 170780788

Effective date: 2022-07-07

MicroVention, Inc.

35 Enterprise Aliso Viejo, CA, 92656 United States of America

Location

497135

MicroVention, Inc.

35 Enterprise Aliso Viejo, CA, 92656 United States of America

Scope

Design, Development, Manufacturing and Distribution of Embolization Prostheses and Accessories, Intravascular Access Devices (Occlusion Balloon Catheters, Guiding and Aspiration Catheters, Microcatheters and Guidewires), Stents, Clot and Foreign Body Retrieval Devices, Liquid Embolic System, Embolic Protection System, Aneurysm Embolization Device, Microspheres, Aspiration Devices and Detachment Controller Units.

499088

MicroVention Costa Rica, S.R.L. Production Site

Zona Franca Coyol Alajuela Costa Rica Manufacturing of Embolization Prostheses and Accessories, Intravascular Access Devices (Occlusion Balloon Catheter, Guiding and Aspiration Catheters, and Microcatheters), Stents, Clot and Foreign Body Retrieval Devices, Embolic Protection System, and Aspiration Devices.





EC Design Examination Certificate

Council Directive 93/42/EEC Annex II Section 4

This is to certify that the manufacturer

MicroVention, Inc.

1311 Valencia Ave. Tustin, CA, 92780 United States of America

that the design of the following device(s)

Scepter C[™] Occlusion Balloon Catheter Scepter XC[™] Occlusion Balloon Catheter Scepter Mini[™] Occlusion Balloon Catheter

is conform to the Essential Requirements of Annex I of the Council Directive 93/42/EEC concerning medical devices.

This EC Design Examination Certificate is only valid in connection with the valid DQS Medizinprodukte GmbH Certificate No. 411133 MR2. Changes to the approved design are subject to further approval by the Notified Body.

Basis of examination: ST18-0008C - Technical Design Dossier For The Scepter Occlusion

Balloon Catheters, April 2019 dated 2019-09-25

Further basis for the examination is referenced in the examination

report and relating documents mentioned below.

Examination report: 411_18e_Report_TFR_Scepter_R2020_V1 dated 2020-02-03

The results of the examination are contained in the above mentioned

report and the relating documents mentioned within.

Certificate registration no. 494215 MRA
Certificate unique ID 170763222
Effective date 2020-02-03
Expiry date 2024-05-26
Frankfurt am Main 2020-02-03

DQS Medizinprodukte GmbH

Sigrid Uhlemann Managing Director Dr. Thomas Feldmann Head of Certification Body

August-Schanz-Straße 21, 60433 Frankfurt am Main, Tel. +49 (0) 69 95427-300, medical.devices@dqs-med.de



DOS (Ne) iz inprodukte GmbH is a Notified Body according to Council Directive 93/42/EEC concerning medical devices with the Identification Number 0297.



EC DECLARATION OF CONFORMITY

RF18-0182, Rev. B

We, MicroVention, Inc., located in Tustin, California, USA, declare according to Directive 93/42/EEC Annex II (excl. Section 4.) under our sole responsibility that the products to which this declaration relates are in conformity with Directive 93/42/EEC and fulfill the Essential Requirements as described in Directive 93/42/EEC Annex I.

Directives:

93/42/EEC Council Directive Concerning Medical Devices

Conformity Assessment Route:

EC Design Examination: Full Quality Assurance:

494215 MRA

411133 MR2

Product	Model Number(s)	Class-Rule	GMDN Code
Scepter XC Occlusion Balloon	BC0411XC		
	BC0410C	_	
Scepter C Occlusion Balloon	BC0415C BC0420C	III – Annex 9, rule 8	32584
	BC0420C	-	
Scepter Mini Occlusion Balloon	BC0210M		

Manufacturer	Notified Body	European Representative
MicroVention, Inc. 1311 Valencia Avenue Tustin, California 92780 USA	DQS Medizinprodukte GmbH Notified Body Number: 0297 D-60433 Frankfurt am Main, Germany	MicroVention Europe 30 bis, rue du Vieil Abreuvoir 78100 Saint-Germain-en-Laye France

Intended Use: For use in the peripheral and neuro vasculature where temporary occlusion is desired. The balloon catheter provides temporary vascular occlusion which is useful in selectively stopping or controlling blood flow. The balloon catheter also offers balloon assisted embolization of intracranial aneurysms. For use in the peripheral vasculature for the infusion of diagnostic agents, such as contrast media, and therapeutic agents such as embolization materials. For neurovascular use for the infusion of diagnostic agents such as contrast media, and therapeutic agents, such as embolization materials, that have been approved or cleared for use in the neurovasculature and are compatible with the inner diameter of the Scepter Occlusion Balloon Catheter.

We herewith declare that the above-mentioned products meet the provisions of the council directive 93/42/EEC for medical devices. All supporting documentation is retained under the premises of the manufacturer.

Irina Kulinets

SVP Regulatory, Quality and Clinical

Lulinet

MicroVention, Inc.

Tustin, CA 92780, USA

Place of Issue

5/30/2019 Date of Issue

Page 1 of 1

Date: 2021.02.23 15:41:35 EET Reason: MoldSign Signature Location: Moldova

Digitally signed by Grabazei Alexandru