

## EC DECLARATION OF CONFORMITY

FD14-0031, Rev. G

We, MicroVention Europe SAR, located in Saint-Germain-en-Laye, France declare according to Directive 93/42/EEC Annex II (incl. 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** 

Annex II, Section 4 - Full Quality System

EC Design Examination: Full Quality Assurance:

517356 MRA Annex II (Section 4)

487703 MR2 Annex II (Excluding Section 4)

Product	Model Number(s)	Class-Rule	Effectivity date	GMDN code
PHIL™ Liquid Embolic System	LEN10250 LEN10250RE LEN10300 LEN10300RE LEN10350 LEN10350RE LEN10LV250 LEN10LV250RE	III – Annex 9, rule 8	2019-03-08	60939

Manufacturer	Notified Body	Production Site
MicroVention Europe SARL 30 bis, rue du Vieil Abreuvoir 78100 Saint-Germain-en-Laye France	DQS Medizinprodukte GmbH Notified Body Number: 0297 D-60433 Frankfurt am Main, Germany	MicroVention, Inc. 1311 Valencia Avenue Tustin, CA 92780, USA Microvention, Inc. 35 Enterprise Aliso Viejo, CA 92656, USA

Intended Use: The PHIL<sup>TM</sup> Liquid Embolic System is intended for use in the embolization of lesions in the peripheral and neurovasculature, including arteriovenous malformations and hypervascular tumors.

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

Sr. VP Regulatory, Clinical, Quality MicroVention Europe SARL

1. Luliack

Place of Issue

Saint-Germain-en-Lave

Date of Issue

Expiry Date: 2022-03-01

Controlled Copy

Ssued Date 4126 19

MASTER

Page 1 of 1