

# EC DECLARATION OF CONFORMITY

RF 19-0097 Rev.B

DC Number: 20-05340

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

## Council Directive 93/42/EEC

### Conformity Assessment Procedure Performed:

<b>EC Design Examination Certificate</b> <input checked="" type="checkbox"/> <u>(Annex II.4)</u>  487703 MRA <b>Certificate Number</b>	<b>EC Full Quality Assurance Certificate</b> <input checked="" type="checkbox"/> <u>(Annex II.3)</u>  487703 MR2 <b>Certificate Number</b>
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Product	Model Number(s)	Class/Rule	GMDN Code
<b>SOFIA® Distal Access Catheter</b>	DA5115ST, DA5125ST, DA6115ST	III – Annex 9, rule 7	58173 (Thrombectomy catheter)
<b>SOFIA® PLUS Catheter</b>	AC6115ST, AC6125ST, AC6131ST, DA6125ST, DA6131ST, DA6135ST		58173 (Thrombectomy catheter)
<b>SOFIA® Flow PLUS Catheter</b>	DA6125ST, DA6131ST		58173 (Thrombectomy catheter)
<b>SOFIA® Flow Catheter</b>	DA5115ST, DA5125ST, AC5115ST, AC5125ST		58173 (Thrombectomy catheter)
<b>SOFIA® EX Catheter</b>	ISC5105ST, ISC5115ST		17846 (Vascular guide catheter, Single Use)

Legal Manufacturer	Production Site(s)	Notified Body
MicroVention Europe SARL 30 bis, rue du Vieil Abreuvour 78100 Saint-Germain-en-Laye France	MicroVention, Inc. 1311 Valencia Avenue Tustin, California 92780 USA  MicroVention, Inc. 35 Enterprise Aliso Viejo, California 92656 USA  MicroVention Costa Rica, S.R.L. Zona Franca Coyol Alajuela, Costa Rica	DQS Medizinprodukte GmbH D-60433 Frankfurt am Main, Germany Notified Body No: 0297

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We herewith declare that the above-mentioned medical device (s) meet the provisions of the council directive 93/42/EEC Medical Device Directive. This declaration is supported by the EC Quality System Certificate (s) according to the provisions of the relevant Annex (es) of above Directive. This declaration applies to all device(s) specified above distributed from the signature date forward.

DocuSigned by:  
  
Signer Name: Irina Kulinets  
Signing Reason: I approve this document  
Signing Time: 11/19/2020 | 6:28:42 PM PST  
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Saint-Germain-en-Laye,  
France

11/19/2020

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Irina Kulinets  
Sr. Vice President, Regulatory  
Affairs, Quality, Clinical Research  
MicroVention Europe SARL

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Place of Issue

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Date of Issue

**Certificate Expiry Date: 2024-05-26**