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СЕРТИФИКАТ

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Benannt durch/Designated by Zentralstelle der Länder für Gesundheitsschutz bei Arzneimitteln und Medizinprodukten ZLG-BS-244,10.08





EC Certificate

Full Quality Assurance System Directive 93/42/EEC on Medical Devices (MDD), Annex II excluding (4) (Devices in Class IIa, IIb or III)

No. G1 106138 0002 Rev. 00

Manufacturer:

Marflow AG

Soodstrasse 57 8134 Adliswil, Zurich SWITZERLAND

Product Category(ies): Class IIb

Double J stent & set Class IIa PCN catheter & set Ureteral catheter Malecot catheter Re-entry malecot catheter Suprapubic catheter Braided shaft catheter Dual lumen catheter Facial dilator Amplatz dilator & set Ureteral dilator & set Ureteral balloon dilator Double J stent & set Mono J stent Endopyelotomy stent Guidewire **IP** Needle Chiba needle Stone basket Perk basket

The Certification Body of TÜV SÜD Product Service GmbH declares that the aforementioned manufacturer has implemented a quality assurance system for design, manufacture and final inspection of the respective devices / device categories in accordance with MDD Annex II. This quality assurance system conforms to the requirements of this Directive and is subject to periodical surveillance. For marketing of class III devices an additional Annex II (4) certificate is mandatory. See also notes overleaf.

Report No.:

IND20190101

Valid from: Valid until: 2020-04-03 2024-05-26



Page 1 of 3 TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123

TÜV SÜD Product Service GmbH / Certification Body - Ridlerstraße 66 - 20339 Munich - Germany



Logalization see reverse side

Official Certification

Seen for authentication of the foregoing signature, acknowledged in our presence by

Ms. <u>Tracey WALTHER</u>, born 28th December 1958, Swiss citizen of Oberentfelden AG, according to her information residing at Brunaustrasse 17, 8002 Zürich, identified by identity card.

Zürich, 8th April 2020 BK no. 1027ff Fee CHF 20.00



NOTARIAT ENGE-ZÜRICH

Andreas Bachmann, Notary Public

	APOSTILLE on de la Haye du 5 octobre 1961)
Land: Schweizerische Eidge Country: Swiss Confederatio Diese öffentliche Urkunde /	in, Canton of Zurich
. ist unterschrieben von	
has been signed by	Andreas Bachmann
in seiner Eigenschaft als	Notary Public
acting in the capacity of	
	tempel/Siegel des (der) - bears the stamp/seal of
Notariat Enge-Zürich Kan	
	Bestätigt / Certified 6. am / the 08.04.2020
. In / at 8090 Zürich / Zurich	
. durch die Staatskanzlei de by the Chancellery of State	s Kantons Zürich e of the Canton of Zurich
unter Nr. / under N° 1	179274/2020
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Legalization see reverse side

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Official Certification

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9. Stempel/Siegel, Stamp/se	al 10. Unterschrift Signature
	S. Overkott
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EC Certificate

Full Quality Assurance System Directive 93/42/EEC on Medical Devices (MDD), Annex II excluding (4) (Devices in class I in sterile conditions, sterilized systems or procedure packs)

No. G1S 106138 0003 Rev. 00

Manufacturer:

Marflow AG

Soodstrasse 57 8134 Adliswil, Zurich SWITZERLAND

Product Category(ies): Class Is Urine bag connector Penile clamp Evacuator IUI catheter without syringe

The Certification Body of TÜV SÜD Product Service GmbH declares that the aforementioned manufacturer has implemented a quality assurance system for manufacture in accordance with MDD Annex II. This quality assurance system covers those aspects of manufacture concerned with securing and maintaining sterile conditions of the respective devices / device categories and conforms to the requirements of this Directive. It is subject to periodical surveillance. See also notes overleaf.

Report No.:

IND20190101

Valid from: Valid until:

2020-04-03 2024-05-26

Date,

2020-04-03

Christoph Dicks Head of Certification/Notified Body

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Page 1 of 2 TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123

TÜV SÜD Product Service GmbH • Certification Body • Ridlerstraße 65 • 80339 Munich • Germany

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