

# EU Quality Management System Certificate

### mdc medical device certification GmbH

Kriegerstr. 6, 70191 Stuttgart, Germany Notified body (identification number 0483)

hereby certifies that the company (SRN: DE-MF-000005600)

## H. + H. Maslanka Chirurgische Instrumente GmbH

Stockacher Straße 172 78532 Tuttlingen Germany

has implemented and applies a quality management system in accordance with Annex IX, Chapter I and III of Regulation (EU) 2017/745 for conformity assessment of the devices listed on the following pages.

An audit by mdc has proven that this quality management system fulfils the following requirements:

# Annex IX - Chapter I (Quality Management System)

of Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices

Surveillance is carried out in accordance with Annex IX, Section 3 of Regulation (EU) 2017/745.

This certificate consists of 2 pages. Details of the devices affected by this certificate as well as further information and conditions are included on the following pages.

 Valid from:
 2023-11-20
 Registration No.
 D1038400076

 Valid until:
 2027-08-17
 Evaluation Report No.
 P23-00107-257670

Stuttgart, 2023-11-20

Head of Notified Body





Devices:
Product: Transseptal needle and transseptal catheter Risk class: III
Product: Flex-Pusher I and II Risk class: III
Product: Unitip-Applicator Risk class: Ila

### Notes:

For the placing on the market of class III and IIb implantable devices (except sutures, staples, dental fillings, dental braces, tooth crowns, screws, wedges, plates, wires, pins, clips and connectors within the meaning of Regulation (EU) 2017/745, Art. 52 (4), 2<sup>nd</sup> paragraph and with the exception of custom-made devices of class III), an EU technical documentation assessment certificate is also required.

The certificate is based on the previous certificate D1038400075 dated 08.05.2023 with the following changes: The product "Transseptal needle" was supplemented by "transseptal catheter".