

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-000010680)

Andreas Hettich GmbH & Co. KG

Föhrenstraße 12 78532 Tuttlingen Germany

has implemented and applies a quality management system in accordance with Annex IX, Chapter I 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:2022-10-25Registration No.D1459300004Valid until:2027-08-25Evaluation Report No.222250

Stuttgart, 2022-10-25

Benannt durch/Designated by Zentralstelle der Länder für Gesundheitsschutz bei Arzneimtteln und Medizinprodukten

BS-MDR-098

Head of Notified Body



Devices:

Product: Centrifuges for separation of blood components for transfusion purposes and for preparatory diagnostics of transfusion blood

Risk class: IIa