

# **EU Quality Management** Certificate



This is to certify that the company



Chemische Fabrik Dr. Weigert GmbH & Co. KG

Mühlenhagen 85 20539 Hamburg Germany

SRN: DE-MF-000005940

has established, implemented and maintains a Quality Management System in accordance with

## Annex IX, Chapter I and III of the Regulation (EU) 2017/745

Conformity Assessment based on a Quality Management System and on Assessment of **Technical Documentation** 

for the device categories and products listed in the Annex of this certificate.

The conformity of the Quality Management System has been verified in an audit and is subject to regular surveillance. Limitations to this certificate are listed in the Annex.

Certificate registration no.	001549 MDR2017Q
Certificate ID	170778872
Effective date	2022-10-13
Expiry date	2027-10-12
Frankfurt am Main,	2022-10-13

### DQS Medizinprodukte GmbH

J. Michael Bothe S. Kuchyn

Sigrid Uhlemann Managing Director

Michael Bothe Head of Certification Body (active medical devices)

nnt durch/Designated by ntralstelle der Länder ür Gesundheitsschutz zneimitteln und edizinprodukten BS-MDR-094

Szymon Kurdyn Head of Certification Body (non-active medical devices)



Accredited Body: DQS Medizinprodukte GmbH, August-Schanz-Str. 21, 60433 Frankfurt am Main DQS Medizinprodukte GmbH is a Notified Body according to Regulation (EU) 2017/745 of the Council concerning medical devices with the Identification Number 0297. The validity of this certificate can only be verified by the QR-code.



## Annex to EU Quality Management Certificate SRN of Manufacturer: DE-MF-000005940 Certificate ID: 170778872

## Authorised Representative of the company:

#### Device categories covered by this certificate:

Device category:	MDN 1211 - Non-active non-implantable devices for disinfecting, cleaning and rinsing
Risk classification:	IIa
Intended purpose:	Products for automated or manual pre-cleaning, cleaning, disinfection of medical devices
Device category:	MDN 1211 - Non-active non-implantable devices for disinfecting, cleaning and rinsing
Risk classification:	IIb
Intended purpose:	Products for automated or manual disinfection of invasive medical devices

**Examinations and tests performed:** 001549\_A209538MED\_01 dated 2022-09-19

#### Further conditions for or limitations to the validity of the certificate:

The manufacturer's quality management system is subject to periodic surveillance in accordance with Annex IX, Chapter 1, Section 3.

#### **Reference to previous certificates:**

Revision	Date of Issue	Certificate-ID	Description of change
n/a	n/a	n/a	n/a