

EU Quality Management Certificate



This is to certify that the company



Miele & Cie. KG

Carl-Miele-Straße 29 33332 Gütersloh Germany

SRN: DE-MF-000005768

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 in accordance with Annex IX, Chapter 1, Section 3. Limitations to this certificate are listed in the Annex.

Devices listed in the Annex may bear the CE marking with the identification number of the Notified Body (0297).

For placing of devices of class III and devices class IIb implantable according to Article 52(4) subparagraph 2 listed in the Annex on the market, an additional certificate according to Annex IX, Chapter II is required.

Certificate registration no. 294819 MDR2017Q

 Certificate ID
 1000215662

 Effective date
 2025-02-27

 Expiry date
 2026-11-03

 Frankfurt am Main,
 2025-02-27



DQS Medizinprodukte GmbH

Heinrich von Mettenheim Managing Director





Annex to EU Quality Management Certificate SRN of Manufacturer: DE-MF-000005768

Certificate ID: 1000215662

Device categories and variants covered by this certificate:

Device category: MDA 0317/A - Active non-implantable devices for cleaning and

disinfection

Product name: PG 8562, PG 8581, PG 8582, PG 8582 CD, PG 8591, PG 8592,

RID-100, RID-200,

PWD 8682, PWD 8692, PWD 8682 CD, PWD 8682 C, RID 101, RID 201,

DS 5010, DS 5011, DS 5020

Risk classification: IIb

Basic-UDI-DI: 4002515GG05MM

Intended purpose: Reprocessable medical devices can be cleaned, rinsed, disinfected and

dried in these Miele washer-disinfectors. These devices can be used in healthcare facilities such as practices, dental practices, hospitals, ambulatory surgical centers or veterinary facilities, depending of the

type

Device category: MDA 0317/A - Active non-implantable devices for cleaning and

disinfection

Product name: ETD PAA Basic, ETD PAA Premium, ETD GA Premium, ETD GA Basic

Risk classification: IIb

Basic-UDI-DI: 4002515ETD6D

Intended purpose: The ETD is an endoscope washer-disinfector intended for automatic

reprocessing for compatible flexible endoscopes. The reprocessing

cycles consists of cleaning, rinsing, disinfection and optional

dehumidifying.

Examinations and tests performed:

294819_A207836MED_01 dated 2021-07-30 420_12d_Bericht_Produktprüfung_Miele_K-Serie_korr dated 19.03.2021 294819_A211325MED_01 dated 13.01.2023 294819_A216149MED dated 2025-02-14

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

n/a

Reference to previous certificates:

Revision 01	Date of Issue 2021-11-04	Certificate-ID 170775965	Description of change Extension Olympus ETD to the models ETD PAA Basic, ETD PAA Premium, ETD GA Premium, ETD GA Basic
02	2023-01-19	170782537	New certificate template
03	2024-05-23	1000169515	(Redesign) of the K series to the models PWD 8682, PWD 8692, PWD 8682 CD, PWD 8682 C, RID 101, RID 201, DS 5010, DS 5011, DS 5020