



# CERTIFICATE



This is to certify that the company



**MEDAGENT**

**MEDAGENT GmbH**

Griesweg 47  
78570 Mühlheim / Donau  
Germany

with the organizational units/sites as listed in the annex  
has implemented and maintains a **Quality Management System**.

Scope:

Design and development of software tools in the field of technical documentation of medical devices, training of medical device manufacturers in the field of development on quality management systems and regulatory requirements for medical devices.

Through an audit, documented in a report, performed by DQS Medizinprodukte GmbH, it was verified that the management system fulfills the requirements of the following standard:

**EN ISO 13485:2016 / A11 : 2021**  
**ISO 13485 : 2016**

Certificate registration no.	547429 MP2021
Certificate unique ID	1000181066
Effective date	2024-06-25
Expiry date	2026-08-06
Frankfurt am Main	2024-06-25



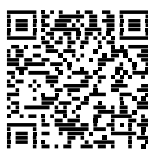
DQS IS A MEMBER OF



**DQS Medizinprodukte GmbH**

Sigrid Uhlemann  
Managing Director

Szymon Kurdyn  
Product Manager



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The validity of the certification can only be verified by the QR-code.



**Annex to certificate**  
**Certificate registration No.: 547429 MP2021**  
**Certificate unique ID: 1000181066**  
**Effective date: 2024-06-25**

## **MEDAGENT GmbH**

Griesweg 47  
78570 Mühlheim / Donau  
Germany

### **Location**

**547708**  
**MEDAGENT GmbH**  
Griesweg 47  
78570 Mühlheim / Donau  
Germany

### **Scope**

Design and development of software tools in the field of technical documentation of medical devices, training of medical device manufacturers in the field of development on quality management systems and regulatory requirements for medical devices.

**31619024**  
**MEDAGENT GmbH**  
Tuttlinger Str. 24  
78579 Neuhausen  
Germany

Design and development of software tools in the field of technical documentation of medical devices, training of medical device manufacturers in the field of development on quality management systems and regulatory requirements for medical devices.