

EU Quality Management System Certificate

We hereby certify the company

Roboscreen GmbH
Hohmannstraße 7
04129 Leipzig
Germany

the introduction and application of a quality management system in accordance with Annex IX, Chapter I and III of Regulation (EU) 2017/746 for conformity assessment.

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

Annex IX – Chapter I (Quality Management System)

of Regulation (EU) 2017/746 of the European Parliament and of the Council of 5 April 2017 on in vitro diagnostic medical devices.

Surveillance is carried out in accordance with Annex IX, Section 3 of Regulation (EU) 2017/746.

This certificate from mdc medical device certification GmbH (Notified Body 0483) consists of 2 pages. Details of the devices covered as well as further information and conditions are contained on the following pages.

Valid from 2026-04-02
Valid until 2029-11-22

Registration No. D1152500053
Report No. P25-00872-359468

Stuttgart, 2026-04-02



Notified Body



Devices:

Devices for the determining the infectious load of a life-threatening disease where monitoring is critical in the process of patient management

Risk class: D

Notes:

For class A devices placed on the market in sterile condition the involvement of mdc is limited to the assessment of the aspects of manufacture concerned with securing and maintaining sterile conditions.

For devices for self-testing and near-patient testing the involvement of mdc additionally refers to the assessment of aspects according to Annex IX, Section 5.1.

For the placing on the market of class D devices an EU technical documentation assessment certificate is also required.