

EU-Quality Management System Certificate

Pursuant to Regulation (EU) 2017/745 on Medical Devices, Annex IX, Chapters I and III

The Notified Body of TÜV NORD CERT GmbH certifies that the manufacturer

Rovers Medical Devices B.V. Lekstraat 10 5347KV Oss The Netherlands

has established, documented and implemented a quality management system in accordance with Article 10, paragraph 9 of Regulation (EU) 2017/745 on medical devices. Details on device categories covered by the quality management system are described on the following page(s). The conformity assessment has been carried out and successfully completed in accordance with Annex IX, Chapters I and III. The result of the assessment, references to investigations carried out, relevant common specifications and test reports are summarised in the report mentioned below. The quality management system assessment was accompanied by the assessment of technical documentation for devices selected on a representative basis. The validity of this certificate is based on the maintenance of the quality management system in accordance with the requirements of the Regulation and its regular surveillance by the Notified Body in accordance with Annex IX, Chapter I, Section 3. The surveillance assessment shall also include an assessment of the technical documentation for the device or devices concerned on the basis of further representative samples.

In addition to the CE marking, the identification number of the Notified Body must be affixed to the devices.

For the placing on the market of class III and Class IIb implantable devices an additional EU technical documentation assessment certificate according to Annex IX, Chapter II is required.

Single Registration Number of the Manufacturer (SRN): Authorised Representative:

The validity of this EU Certificate depends on conditions

and / or is limited to the following:

NL-MF-000001553

N/A

List of Products, Risk Classification and Details:

Certificate history:

see Section 2

see Section 3

Reg.-No.: 44 911 220665

Certification decision report No.: 3535 9389

Edition: 1

Issue date: 2024-05-17 First issued: 2024-05-17 Valid until: 2029-05-16

Essen, 2024-05-17

B. Hoy

TÜV NORD CERT GmbH is a Notified Body with identification number 0044

TÜV NORD CERT GmbH Am TÜV 1, 45307 Essen tuev-nord-cert.de medical@tuev-nord.de TÜV®





EU-Quality Management System Certificate

Pursuant to Regulation (EU) 2017/745 on Medical Devices, Annex IX, Chapters I and III

Reg. No. 44 911 220665 Section 2, List of Products

Class I, sterile

Product name	Category of device (MDx)	Technical documentation assessment report number
Rovers® Anex® Brush	MDN 1202,	3534 3444
Rovers® Cervex-Brush®	MDS 1005	
Rovers® Cervex-Brush® Combi		
Rovers® EndoCervex-Brush®		
Rovers® EndoCervex-Brush®-S		
Rovers® Orcellex® Brush		
Rovers® Viba-Brush®		
Rovers® Evalyn® Brush		

For class is devices placed on the market in a sterile condition, the involvement of the notified body in the conformity assessment procedure is limited to those aspects related to the manufacture, securing and maintenance of sterile conditions.





EU-Quality Management System Certificate

Pursuant to Regulation (EU) 2017/745 on Medical Devices, Annex IX, Chapters I and III

Reg. No. 44 911 220665 Section 3, Certificate History

Certificate History

Edition	Date	Action leading to revision	Certification decision report number
01	2024-05-17	Initial issuance	3535 9389

