

EU Quality Management System Certificate

We hereby certify the company

Risa GmbH Industriestraße 7 78234 Engen Germany

the introduction and application of a quality management system in accordance with Annex IX, Chapter I and III of Regulation (EU) 2017/745 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) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices.

Surveillance is carried out in accordance with Annex IX, Section 3 of Regulation (EU) 2017/745.

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

Valid from 2024-07-11 Valid until 2029-06-17 Registration No. D1084100033 Report No. P23-00692-268700

Stuttgart, 2024-07-11

Notified Body



Risa GmbH | SRN: DE-MF-000011054

Devices:

Risk class: Ila

Bone Drills and Reamers

Registration No. D1084100033