



EC Design Examination Certificate

Council Directive 93/42/EEC Annex II Section 4

This is to certify that the manufacturer



Gebrüder Martin GmbH & Co. KG

KLS Martin Platz 1 78532 Tuttlingen Germany

that the design of the following device(s)

HBS 2 Resorb Mg

is conform to the Essential Requirements of Annex I of the Council Directive 93/42/EEC concerning medical devices.

This EC Design Examination Certificate is only valid in connection with the valid DQS Medizinprodukte GmbH Certificate No. 210299 MR2. Changes to the approved design are subject to further approval by the Notified Body.

Basis of examination: 13-014-01_STED_001-pdf dated 2019-10-25

13-014-01_STED_001.pdf dated 2021-11-30

Further basis for the examination is referenced in the examination

report and relating documents mentioned below.

Examination report: 411_18e_Report_TFR_CHS_V2.docx dated 2020-06-10

411_18e_Report_TFR_CHS_V3.docx dated 2021-04-20

The results of the examination are contained in the above mentioned

report and the relating documents mentioned within.

Certificate registration no. 548582 MRA
Certificate unique ID 170774687
Effective date 2021-04-20
Expiry date 2024-05-26
Frankfurt am Main 2021-04-20

DQS Medizinprodukte GmbH

Sigrid Uhlemann Managing Director Dr. Thomas Feldmann Head of Certification Body

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Annex to certificate

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Device Family: Devices:

