

3EC International a.s., Hraničná 18, 821 05 Bratislava, Slovak Republic Notified body No. 2265

EU QUALITY MANAGEMENT SYSTEM CERTIFICATE No. 2022-MDR/QS-005

Healthium Medtech Limited

Registered Place of Business: No. 472 D, 13th Cross, 4th Phase, Peenya Industrial Area,

Bangalore, Karnataka - 560 058, India

Manufacturing Site 01: No. 472 D, 13th Cross, 4th Phase, Peenya Industrial Area, Bangalore,

Karnataka - 560 058, India

Manufacturing Site 02: Plot No: 1605, Portia Road, Sri City, Tirupati District,

Andhra Pradesh - 517 646, India SRN No.: IN-MF-000008421

Name and address of the Authorized representative:

MED DEVICES LIFESCIENCES B.V., Keizersgracht 482, 1017 EG Amsterdam, The Netherlands

This EU Quality Management System Certificate issued in accordance with the Regulation (EU) 2017/745 of the European Parliament and of the Council on medical devices as amended confirms, that quality management system of medical device.

BRAIDED AND COATED POLYGLYCOLIC ACID

STERILE SYNTHETIC ABSORBABLE SURGICAL SUTURE

(for detailed list refer to Annex I) Intended purpose: Annex II

MD class III

meets the requirements on quality management system according to the Chapter I Annex IX of the Regulation (EU) 2017/745 of the European Parliament and of the Council on medical devices as amended.

Conditions for or limitations to the validity of the certificate: N/A

Validity of the certificate is conditional upon positive results of regular surveillance audits.

Notified body No. 2265 has performed assessment of the quality management system of the abovementioned medical device and found that it meets the requirements stated above. The outcome of the assessment of the quality management system of the abovementioned medical device is stated in the MD Technical Documentation Assessment Report No. MDR020_2021 from August 4. 2022, MD Clinical Evaluation Report No. MDR020_2021 from August 4. 2022 and MD Audit Report No. SK-0643-22 from August 4. 2022. Information on all examinations and tests performed is stated in the abovementioned reports and is available on request.

This EU Quality Management System Certificate applies only to the quality management system of the abovementioned medical device. For the placing on the market of the MDs which this certificate covers, the EU Technical Documentation Assessment Certificate issued in accordance with the Regulation (EU) 2017/745 on medical devices as amended is required. The certificate validity is conditional upon fulfilment of relevant legal requirements by the manufacturer.

Valid from: October 18, 2022 Valid until: August 15, 2027

First issue: August 15, 2022

Revision: 01 History: Annex III BEC International

3EC International a.s. Katarina Tomin Srdošová, PhD. Director of NB2265

In Bratislava, Slovakia, October 18, 2022-



ANNEX I TO EU QUALITY MANAGEMENT SYSTEM CERTIFICATE No. 2022-MDR/QS-005

issued for the company

Healthium Medtech Limited

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List of medical devices covered by the EU Quality Management System Certificate:

BRAIDED AND COATED POLYGLYCOLIC ACID STERILE SYNTHETIC ABSORBABLE SURGICAL SUTURE				
Brand Name	USP sizes	EP metric size		
TRUGLYDE, ANNCRYL, I GLYDE, LINX PGA, POLYGLYCOLIC ACID, U-GLYDE, PGA RESORBA, SURGISUT TRUGLYDE, B-GLYDE, TRUVET POLYGLYCOLIC ACID, IM-GLYDE, ESORB, UNODENT PGA, POGAL, N-CARE PGA, ALAN GLYDE	9-0. 8-0. 7-0. 6-0. 5-0. 4-0. 3-0. 2-0. 0. f. 2	0.3, 0.4, 0.5, 0.7, 1, 1.5, 2, 3, 3.5, 4, 5		

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Katarina Tomin Srdošová, PhD. Director of NB2265

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* Suture is supplied with or without needles.



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Intended purpose of medical devices covered by the EU Quality Management System Certificate:

Braided and coated Polyglycolic acid sutures is intended for use in general soft issue approximation and/or ligation, including use in plastic and ophthalmic procedures, but not for use in cardiovascular and neurological tissues.

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katarina Tomin Srdošová, PhD Director of NB2265

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ANNEX III TO EU QUALITY MANAGEMENT SYSTEM CERTIFICATE No. 2022-MDR/QS-005

issued for the company

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Certificate history:

Revision	EU QMS Certificate reference	Date of issue	Application for Conformity Assessment of MD number	Description
00	2022-MDR/QS-005	15.8.2022	MDR020_2021	Initially granted certification
01	2022-MDR/QS-005	18.10.2022	MDR020 2021	Correction of typo mistake in the manufacturing site 02 address

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