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EU-Declaration of Conformity for Medical Device Class I

Hamburg, 2022-12-05

Object(s) of the declaration:

Bodedex forte

Bodedex forte		
Pack size	Article number BODE	Article number HARTMANN
21	973762	980244
51	973761	980243
	973769	980250

We herewith declare under our sole responsibility that the medical devices listed above, first placed on the market by BODE Chemie GmbH, comply with the applicable provisions, in particular, the

• General Safety and Performance Requirements of Regulation (EU) 2017/745 of the European Parliament and of the Council of 5. April 2017 on medical devices.

The objects of the declaration have been identified as medical devices in risk class I according to classification rule 1 in Annex VIII of Regulation (EU) 2017/745. The conformity assessment procedure according to Article 52 (7) has been performed and the Technical Documentation is kept available.

Intended Purpose: Cleaning of invasive and non-invasive medical devices.

Basic UDI-DI: 40316783776ME Single Registration Number: DE-MF-000005851 Certificate No. 0523GB448210329A

BODE Chemie GmbH

Dr. Henning Mallwitz Director Research & Development

0 6. DEZ. 2022

Dr. Ralf Meier Head of Quality Assurance

Valid until: 2024-12-05



HARTMANN SCIENCE CENTER Research for infection protection. BODE Chemie GmbH Commercial Register Hamburg HRB 108924 Page 1 of 1 Managing Director: Alexander Schwieger HypoVereinsbank IBAN: DE83 6002 0290 0611 0224 33 BIC: HYVEDEMM473 · VAT-ID: DE118115768