BODE Chemie GmbH

A company of the HARTMANN GROUP

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EC-Declaration of Conformity for Medical Device Class Ila

Hamburg, 2021-11-16

We herewith declare, that

Object of the declaration:

Mikrobac forte

| Pack size | Article number BODE | Article number HARTMANN |
|--------------------|----------------------------|----------------------------|
| 250 x 20 ml sachet | 975392 | 980434 |
| 5 I canister | 975395 981179 973218 | 980435 981179 980184 |
| 200 l drum | 975397 | 980437 |
| 640 I container | 975398 | 980438 |

which is manufactured and/or placed on the market by BODE Chemie GmbH, meet the applicable provisions, especially the essential health and safety requirements of the following EC-Directive:

Council Directive 93/42/EEC of 14th June, 1993

The required conformity assessment procedure according to Annex II excluding (4) has been performed and the technical documentation is kept available.

This EC-Declaration of Conformity is issued under the sole responsibility of the BODE Chemie GmbH.

The conformity assessment procedure is under the supervision of the Notified Body:

MEDCERT Zertifizierungs- und Prüfungsgesellschaft für die Medizin GmbH, Pilatuspool 2, 20355 Hamburg, Germany

Identification No. 0482

Medical Device: Class IIa acc. to rule 15 (acc. to Annex IX of the directive)

BODE Chemie GmbH

Dr. Henning Mallwitz

Director Research & Development

This document is valid until: 2023-02-08

André Maack

Head of Quality Assurance