## **BODE Chemie GmbH**

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## **EU-Declaration of Conformity for Medical Device Class IIb**

Hamburg, 2023-11-07

Object of the declaration:

## Mikrobac Virucidal Tissues

| Mikrobac Virucidal Tissues |                     |                         |
|----------------------------|---------------------|-------------------------|
| Pack size                  | Article number BODE | Article number HARTMANN |
| Flowpack 80 Tissues        | 981531              | 981531                  |

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 IIb according to classification rule 1 and rule 16 in Annex VIII of Regulation (EU) 2017/745.

The conformity assessment procedure according to Article 52 (4) and Annex IX has been performed and the Technical Documentation is kept available.

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 Certificate No. 0523GB448210329A

(High-Level) Intended Purpose:

Disinfection of non-invasive and invasive medical devices

Basic UDI-DI: 40316783779ML

Single Registration Number: DE-MF-000005851

**BODE Chemie GmbH** 

Britta Glogau

Director Research & Development

Valid until: 2025-11-07

Raphael Bohner Head of Quality