POHL BOSKAMP



Declaration of Conformity

We, the company

G. Pohl-Boskamp GmbH & Co. KG

Kieler Strasse 11

25551 Hohenlockstedt, Germany

Single Registration Number: DE-MF-000010001

declare on our own responsibility, that the risk class IIa medical device (rule 21, dash 3 of annex VIII of the regulation mentioned below)

GeloRevoice® Throat Lozenges with

Intended Purpose: GeloRevoice[®] Throat Lozenges provide protection of the mouth and throat lining (mucous membranes). They are used to relieve throat maladies such as hoarseness, throat irritation, tickly throat and dry mucous membranes, which lead to a sore throat or an unproductive, dry cough.

Product code: 577

Basic-UDI-DI: 40291250012G

which is subject of this declaration, complies with the requirements of Annex I

and this Declaration of Conformity complies with the Annex IV

of the REGULATION (EU) 2017/745 of 5th of April, 2017 on medical devices (MDR).

The Notified Body

DNV MEDCERT GmbH

Pilatuspool 2

20355 Hamburg, Germany

with the Identification Number 0482 was involved in the conformity assessment according to Annex IX of the REGULATION (EU) 2017/745 and has issued the certificate no.

3253GB448250403 valid until 06 December 2026.

This Declaration of Conformity is valid until 06 December 2026.

Boskarnip GmbA

Hohenlockstedt, 17 April 2025

POHL BOSKAMP

Ines Rowedder

Person responsible for regulatory compliance

(PRRC) acc. to Article 15 MDR

Lea Dornseiff

Advisor Medical Device Registration

RECYCLING