

Manufacturer: Beurer GmbH (see address in footer)

SRN: DE-MF-000005422

Product category: CPR devices

Product type: RH 112

The product specified above is in conformity with the following specifications.

(EU) 2017/745 Medical device regulation (MDR)

Basic-UDI-DI: 4211125RH112EY

Classification/applied rule(s): Class I/rule 13

Conformity assessment procedure: not applicable for class I devices

Certificate no. and validity: D1311700056, valid to 2026-11-29

2011/65/EU Restrictions of the use of certain hazardous substances in electrical and electronic equipment (RoHS)

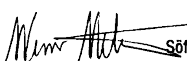
EN IEC 63000:2018

This declaration of conformity is issued under the sole responsibility of the manufacturer.

Signed for and on behalf of: Beurer GmbH

Place, date of issue: Ulm, 2023-11-30

Name, function, signature, stamp: Werner Meternek, Director Quality Management & Regulatory Affairs

ppa.  **Beurer GmbH**  
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