

File

Title EC Declaration of Conformity

PO 40_EC DoC_Beurer_20200508

Manufacturer:

Beurer GmbH (see address in footer)

Product category:

Pulse oximeter

Product type:

PO 40

The product specified above is in conformity with the provisions of the following Union legislation and harmonized standards.

93/42/EEC

Medical device directive (MDD)

UMDNS code and name:

17-148 Oximeter, Pulse

Classification/applied rule(s):

Class Ila/rule 10

Conformity assessment

Annex II - excluding section 4

procedure:

The notified body mdc medical device certification GmbH, located at Kriegerstr. 6, 70191 Stuttgart, Germany, identification number 0483, issued in the course of the mentioned conformity assessment procedure the following certificate:

Certificate no. and validity:

D1311700043, valid to 2024-05-26

IEC 60601-1: 2012 EN 60601-1-2:2015 ISO 80601-2-61:2011

2011/65/EU

Restrictions of the use of certain hazardous substances in electrical and electronic

equipment (ROHS)

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, 08.05.2020

Name, function, signature,

stamp:

Daniel Kämmerer, Team Leader RA

Beurer GmbH
(Söflinger Straße 218 • 89077 Ulm