

# EU Declaration of Conformity

Certificate No.: MLIFR2024012502

*Manufacturer:*

**Microlife Corporation**  
**9F, No. 431, RuiGuang Road, NeiHu,**  
**Taipei, 114, Taiwan, R.O.C.**  
 Single Registration Number(SRN): TW-MF-000010688

*whose single Authorized Representative:*

**Microlife UAB**  
**P. Lukšio g. 32**  
**08222 Vilnius, Lithuania**  
 Single Registration Number(SRN): LT-AR-000011673

We, the manufacturer, herewith declare that the products

**Digital Infrared Thermometer Series**  
**Basic UDI-DI: 4719003IRSR**

**Class: IIa**  
**Trade Name: Microlife**

Customers type no.:	Manufacturers type no.:	EMDN Code	GMDN Code
IR 150	IR 150 (ERP IR1DF1-1)	V0301010202	17887
IR 210	IR 210 (ERP IR1DN1)	V0301010202	17887
NC 150	NC 150 (ERP FR1MF1)	V0301010202	17888
NC 200	NC 200 (ERP FR1DG1)	V0301010202	17888

**Intended purpose:** The devices are intended to measure body temperature.

meet the provisions of Medical Device Regulation (EU) 2017/745 which apply to them.

The medical devices have been assigned to class IIa according to Annex VIII Rule 10 of the Medical Device Regulation (EU) 2017/745.

It bears the mark



The products concerned have been designed and manufactured under a quality management system according to Annex IX of Medical Device Regulation (EU) 2017/745.

Compliance of the designated products with the Medical Device Regulation (EU) 2017/745 have been assessed and certified by the Notified Body

**SGS Belgium NV**  
**Noorderlaan 87 2030 Antwerp Belgium**  
 Certificate No.: TW23/00000625  
 Validity from: 2023-12-21 Expiry date: 2028-8-24

Following the procedure relating to the EU Declaration of Conformity set out in Annex IV of Medical Device Regulation (EU) 2017/745, and in conformity to the following standards or other normative documents:

- EN 60601-1:2006+A1:2013+AC:2014+A12:2014+A2:2021(IEC 60601-1:2005+AMD1:2012+AMD2:2020) or EN 60601-1:2006+A1:2013+AC: 2014 (IEC 60601-1:2005+CORR.1:2006+CORR.2:2007+AM1:2012(or IEC60601-1:2012))
- EN 60601-1-2:2015+A1:2021(IEC 60601-1-2:2014+AMD1:2020) or EN 60601-1-2:2015 (IEC 60601-1-2:2014)
- EN 60601-1-11:2015+A1:2021(IEC 60601-1-11:2015+AMD1:2020)
- EN 60601-1-6:2010+A1:2015+A2:2021(IEC 60601-1-6:2010+AMD1:2013+AMD2:2020)
- EN 62366-1:2015+AC:2015+AC:2016+A1:2020(IEC 62366-1:2015+AMD1:2020)
- EN 62304:2006+A1:2015(IEC 62304:2006+A1:2015)
- EN ISO 80601-2-56:2017+A1:2020(ISO 80601-2-56:2017+A1:2018)
- EN ISO 10993-1:2020(ISO 10993-1:2018)
- EN ISO 10993-5:2009(ISO 10993-5:2009)
- EN ISO 10993-10:2023(ISO10993-10:2021)
- EN ISO 10993-12:2021(ISO 10993-12:2021)
- EN ISO 14971: 2019+A11:2021(ISO 14971: 2019)
- EN ISO 15223-1:2021(ISO 15223-1:2021)
- EN ISO 13485:2016+A11:2021
- ISO 20417:2021
- ISO 17664-2:2021
- MEDDEV 2.7/1 revision 4
- 2011/65/EU amended by M81 (2023/171) and corrected by C2

EC/1907/2006 amended by M74 (2023/1464) and corrected by C9

The above mentioned declaration of conformity is issued under the sole responsibility of Microlife Corporation, and this declaration is valid until Aug 24, 2028.

Place and Date of issue: Taipei, 25 Jan 2024

A handwritten signature in black ink, appearing to read 'Ariel Wang', written in a cursive style.

*Ariel Wang,*

*Management Representative, PRRC*