

**1. Manufacturer and or Authorised Representatives Details****Manufacturer**

Name: Leica Microsystems CMS GmbH  
Address: Ernst-Leitz-Straße 17-37,  
35578 Wetzlar, Germany

**UKRP**

Name: Leica Microsystems (UK) Limited  
Address: Larch House, Woodlands Business Park,  
Milton Keynes,  
England, United Kingdom, MK14 6FG

**2. This declaration is used under the sole responsibility of the manufacturer**

We hereby declare that the device described below, both in its basic design and construction and in the version marked by us, conforms to the relevant safety- and health related requirements of the appropriate UK legislations.  
This declaration shall cease to be valid if modifications are made to the device without our approval.

**3. Object of the declaration**

Product: Microscope  
Model: DM1000 LED  
Order no.: 11 888 842  
Product class: A (according to rule 5 Annex VIII of EU directive 2017/746)  
GMDN Code: 15132 (Microscopic blood / cell examination Instruments IVD)  
Basic UDI-DI: 42519697DM1000AF (Basic Unique Device Identification)

**4. First date of UKCA compliance**

Date: January 11<sup>th</sup> 2023

**5. The products listed are in compliance with the relevant statutory requirements detailed**

Medical Device Regulations 2002 (SI 2002 No 618) Part IV In Vitro Diagnostic Medical Devices

The Restriction of the Use of Certain Hazardous Substances in Electrical and Electronic Equipment Regulations 2012

**6. References to the designated standards used to which conformity is declared**

|                          |                         |
|--------------------------|-------------------------|
| BS EN 61010-1+A1:2017    | EN 61010-1:2010+A1:2019 |
| BS EN 61010-2-101:2017   | EN 61010-2-101: 2017    |
| BS EN IEC 61326-1:2021   | EN 61326-1:2013         |
| BS EN IEC 61326-2-6:2021 | EN 61326-2-6:2013       |
| BS EN 62471:2009         | EN 62471:2008           |
| BS EN IEC 63000:2018     | EN IEC 63000:2018       |

**7. Details of Approved Body**

n.a.

**8. Signed on behalf of the manufacturer by**

Name: Steffen Laabs  
Job Title: RA/QA Director

Date: 11.01.2023  
Signed: 