

## EU DECLARATION OF CONFORMITY

We, **Leica Biosystems Nussloch GmbH, Heidelberger Str. 17-19, 69226 Nussloch, Germany**

hereby declare under our sole responsibility that the medical device

Product and Trade name	<b>Leica CM3050 S</b>
Product	Cryostat Microtome
Risk Class	A
Basic UDI-DI	010404918804709R
Single Registration Number	DE-MF-000021943
Product description	A precision cutting instrument contained within a temperature-controlled cabinet (i.e., a cryostat) intended to be used for the sectioning of rapidly-frozen tissue specimens without prior fixation to expedite subsequent in vitro diagnostic analysis of the tissue specimen.

meets the provision European legislation:

- Regulation (EU) 2017/746 of the European Parliament and of the Council of 5 April 2017 on in vitro diagnostic medical devices (OJ L 117, 5.5.2017, p. 176–332). The procedure according to Annex II and Annex III of the above-mentioned regulation has been followed.

EN 61010-2-101:2017  
EN ISO 14971:2019  
EN 61326-2-6:2013

- Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment (OJ L 174, 1.7.2011, p. 88–110)
- Commission Delegated Directive (EU) 2015/863 of 31 March 2015 amending Annex II to Directive 2011/65/EU of the European Parliament and of the Council as regards the list of restricted substances (OJ L 137, 4.6.2015, p. 10–12)

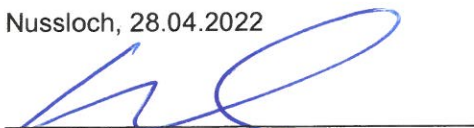
EN IEC 63000:2018

Quality Management System: Certified according to EN ISO 13485:2016 and ISO 9001:2015

Manufacturing sites: Leica Biosystems Nussloch GmbH, Heidelberger Str. 17-19,  
69226 Nussloch, Germany

This declaration is effective for products placed on the market as of the date of issue. Any modification of the device not authorized by Leica Biosystems will invalidate this declaration.

Nussloch, 28.04.2022



Andreas Eich  
Senior Director CH Nussloch



Robert Gropp  
RA/QA Director

Rev. B