

UKCA Declaration of Conformity for LumiraDx INR Test Strips (S-RA-TEC-0002)

Document Number:	S-RA-REP-00462	Revision:	1
Information Classification		Public	



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UK Declaration of Conformity for LumiraDx INR Test Strips (EN)

We, the legal manufacturer, as stated below, herby declares under our sole responsibility,

Legal Manufacturer:	LumiraDx UK Ltd.	
Address:	Dumyat Business Park, Alloa FK10 2PB, United Kingdom	

that the identified products to which this declaration relates,

Identification		
LumiraDx INR Test Strips		
L0030001nnxxx (where nn represents two digits corresponding to language variants, and where xxx represents three digits corresponding to number of units the catalogue number contains).		
55983 – Prothrombin time (PT), IVD, Kit, Clotting		
General IVD		
The Medical Devices Regulations 2002 (SI 2002 No 618, as amended) (UK MDR 2002) through Annex III (excluding Section 6) of 98/79/EC		

is in conformity with the following UK Regulations and Directives as transposed into the national laws of the member states:

Directives and Regulations	
SI 2002 No 618 (As Amended) Medical Devices Regulations 2002	

This declaration is also supported by the following applied standards:

Standards/Common Specifications		
EN ISO 13485:2016	Medical Devices – Quality Management Systems – Requirements for Regulatory Purposes	
EN ISO 14971:2019	Medical Devices – Applications of Risk Management to Medical Devices	

Full responsibility and compliance of the product is signed for and on the behalf of Manufacturer LumiraDx UK Limited:

Name	David Scott	Position:	Chief Technology Officer
Signature		Date:	
72,	SOH.		24 May 2022

With approval of this Declaration of Conformity, we hereby affix the UKCA Mark to the product.

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