



## UKCA Declaration of Conformity for LumiraDx SARS-CoV-2 & Flu A/B and Quality Controls (S- RA-TEC-0012)

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## UK Declaration of Conformity for LumiraDx SARS-CoV-2 & Flu A/B (EN) and LumiraDx SARS-CoV-2 & Flu A/B Quality Controls (EN)

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

<b>Legal Manufacturer:</b>	LumiraDx UK Ltd.
<b>Address:</b>	Dumyat Business Park Alloa FK10 2PB United Kingdom

that the identified products to which this declaration relates,

Identification	
<b>Product Name</b>	LumiraDx SARS-CoV-2 & Flu A/B
<b>Catalogue Numbers</b>	L0190001nnxxx (where nn represents two digits corresponding to language variants, and where xxx represents three digits corresponding to number of units the catalogue number contains).
<b>GMDN Code</b>	48243 - Multiple respiratory virus antigen IVD, kit, fluorescent immunoassay
<b>Classification</b>	General IVD
<b>Conformity Assessment Route</b>	The Medical Devices Regulations 2002 (SI 2002 No 618, as amended) (UK MDR 2002) through Annex III (excluding Section 6) of 98/79/EC
<b>Product Name</b>	LumiraDx SARS-CoV-2 & Flu A/B Quality Controls
<b>Catalogue Numbers</b>	L0190801nnxxx (where nn represents two digits corresponding to language variants, and where xxx represents three digits corresponding to number of units the catalogue number contains).
<b>GMDN Code</b>	48247 - Multiple respiratory virus antigen IVD, control
<b>Classification</b>	General IVD
<b>Conformity Assessment Route</b>	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:		
Original Declaration of Conformity signed (S-RA-REP-00358 Rev 1) by David Scott	22 December 2021		
	10 Nov 2022		

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