

## IVD Technical File Declaration of Conformity for LumiraDx SARS-CoV-2 Ag Test Strips (S-RA-TEC-0007)

Document Number:	S-RA-REP-00127	Revision:	2
Information Classification		Public	



## **Contents**

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## EC Declaration of Conformity for LumiraDx SARS-CoV-2 Ag 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
EC Authorized Representative:	LumiraDx AB
	Västra Vägen
	5A 169 61
	Solna
	Sweden

that the identified products to which this declaration relates,

Identification			
Product Name	LumiraDx SARS-CoV-2 Ag Test Strips		
Catalogue Numbers (without swabs)	L0160001nnxxx (where nn represents two digits corresponding to language variants, and where xxx represents three digits corresponding to number of units the catalogue number contains).		
Catalogue Numbers (with swabs)	L0160006nnxxx (where nn represents two digits corresponding to language variants, and where xxx represents three digits corresponding to number of units the catalogue number contains).		
GMDN Code	66464, SARS-CoV-2 antigen IVD, kit, rapid microfluidic immunoassay, clinical		
Classification	General IVD		
Conformity Assessment Route	Annex III (excluding Section 6) of 98/79/EC		

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

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Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on in vitro diagnostic medical devices

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## Signed for and on the behalf of Manufacturer LumiraDx UK Limited:

Name	David Scott	Position:	Chief Technology Officer
Signature		Date:	
Original Declaration Rev 1) by David Sco	n of Conformity signed (S-RA-REP-00127 tt		26 August 2020
DSCAD.		10 Nov 2022	

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

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