

# EC Declaration of Conformity

**Manufacturer:**

**Name:** Triplex International Biosciences (China) Co., LTD.  
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**Whose Authorized Representative:**

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We, Triplex International Biosciences (China) Co., LTD.(Manufacturer), here declare that the below mentioned medical device meets the provisions of Directive 98/79/EC which apply to them. The declaration of conformity is exclusively under the responsibility of Triplex International Biosciences (China) Co., LTD.(Manufacturer).

<b>Product Name</b>	SARS-CoV-2 Neutralizing antibody Rapid Test Kit				
<b>Intended Use</b>	This SARS-CoV-2 Neutralizing antibody Rapid Test Kit is only used for rapid in vitro qualitative assessment of SARS-CoV-2 neutralizing antibody in human serum/plasma/whole blood(Venipuncture and Fingerstick). For professional In Vitro Diagnostic Use Only.				
<b>PackSize/REF/ Barcodes</b>	<i>PackSize:</i> <i>REF:</i> <i>Barcodes:</i>	1 test/kit C011911 6950917930171	5 tests/kit C011912 6950917930188	10 tests/kit C011913 6950917930195	25 tests/ kit C011914 6950917930201
<b>Classification</b>	Others				

**Conformity Assessment Route:** IVDD 98/79/EC Annex III(excluding Annex III.6).

**Applicable Standards:**

**QMS:**

EN ISO 13485:2016

**Risk Management:**

ISO 14971:2019

**Product Standards:**

EN ISO 18113-1:2011    EN ISO 18113-2:2011

EN 13612:2002    EN 13641:2002    EN 62366-1:2015

ISO 15223-1:2016    ISO 23640:2015    ISO 10993



**Valid until:** May 24, 2022

<b>Name Of Authorized Signatory</b>	Jin Li (李劲)
<b>Position Held In The Company</b>	General Manager
<b>Signature</b>	
<b>Date</b>	July 5, 2021
<b>Place</b>	Xiamen, China.
<b>Seal (Manufacturer)</b>	