

# EU TECHNICAL DOCUMENTATION ASSESSMENT CERTIFICATE

Manufacturer:

ANHUI DEEPBLUE MEDICAL TECHNOLOGY CO., LTD.

No. 777 Jimingshan Road, High-Tech Development Zone, 230088

Hefei, Anhui PEOPLE'S REPUBLIC OF CHINA

Single registration number:

CN-MF-000018785

Authorized representative:

Mega Eurostar Sp. z o. o.

Obrzeżna, 5 lok. XIP/1 02-691 Warsaw

Poland

Single registration number:

PL-AR-000042730

Notified Body Sertio Oy declares that the requirements of Annex IX, Chapter II of the

## REGULATION (EU) 2017/746 on In Vitro Diagnostic Devices

have been met for the products listed in this certificate.

The above mentioned manufacturer has established and maintains a technical documentation defined by Annex IX chapter II. In addition to this certificate an EU Quality Management System certificate is required before placing the listed product on the market.

Certificate validity is subject to manufacturer fulfilling the obligations arising from the Annex IX of the aforementioned regulation. Validity of the certificate is subject to following the General terms of Business by Sertio Oy and Terms of conformity assessment of IVD medical devices.

Certificate number

EU-TDA-FI-20642-800030-2025-1

Issue date

20.05.2025

Valid from

20.05.2025

Expiry date

21.03.2030

Mikko Soikkeli

**Sertio Oy** 

Biokatu 10, 33520 Tampere, Finland





#### **PRODUCTS**

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#### Class C for self testing

**IVR 0503** 

Devices intended to be used to detect the presence of, or exposure to an infectious

agent including sexually transmitted agents

W0105099099 Virology - RT & POC - other

Product name: SARS-CoV-2 & Influenza A+B & RSV& ADV Antigen Combo Test Kit(Colloidal Gold)

Model:

CFRA1ST-X Basic-UDI-DI: 69520627J034LD

Intended use: This product is used for the qualitative detection of SARS-CoV-2, influenza A,

influenza B, respiratory syncytial virus (RSV) and adenovirus antigen in human nasal swab specimens. It is a non-automated rapid test method for infection. This test is authorized for non-prescription home use with self collected anterior nasal (nares) swab samples from individuals. Individuals who test positive should seek follow up care with their physician or healthcare provider as additional testing may be

necessary. Users under the age of 15 should complete the test with supervision of an

adult. Both symptomatic and asymptomatic infections can be tested.

**IVR 0503** 

Devices intended to be used to detect the presence of, or exposure to an infectious

agent including sexually transmitted agents

W0105099099 Virology - RT & POC - other

Product name: Influenza A+B & COVID-19 (SARS-CoV-2) Antigen Test Kit (Colloidal Gold)

Model:

COVAg+FluAB1NST-X

Basic-UDI-DI: 69520627J017LD

Intended use: This product is used for the qualitative detection of SARS-CoV-2, influenza A and influenza B in human nasal swab specimens. It is a non-automated rapid test method for infection. This test is authorized for non-prescription home use with self-collected anterior nasal (nares) swab samples from individuals. Individuals who test positive should seek follow up care with their physician or healthcare provider as additional testing may be necessary. Users under the age of 15 should complete the test with supervision of an adult. Both symptomatic and asymptomatic infections can be

tested.



### **Certificate history**

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Version	Date issued	Description
1	21.03.2025	Initial certification
2	20.5.2025	Addition of device: Influenza A+B & COVID-19 (SARS-CoV-2) Antigen Test Kit (Colloidal Gold), Basic UDI-DI: 69520627J017LD

