



CERTIFICATE

EC Certificate No. 1434-IVDD-035/2022

**EC Design-examination
Directive 98/79/EC concerning
in vitro diagnostic medical devices**

Polish Centre for Testing and Certification certifies
that manufactured by:

Hangzhou AllTest Biotech Co., Ltd
**550#, Yin Hai Street, Hangzhou Economic & Technological
Development Area, Hangzhou, 310018, P.R. China**

**in vitro diagnostic medical devices
for self-testing**

SARS-COV-2 Antigen Rapid Test (Nasal Swab)

The list of medical devices covered by this certificate is provided in the annex 1

in terms of design documentation, comply with requirements
of Annex III (Section 6) to Directive 98/79/EC (as amended)
implemented into Polish law,
as evidenced by the audit conducted by the PCBC

Validity of the Certificate: from 16.03.2022 to 27.05.2025

The date of issue of the Certificate: 16.03.2022

The date of the first issue of the Certificate: 05.07.2021



Issued under the Contract No. MD-34/2021
Application No: 048/2021
Certificate bears the authorized person signature.
Warsaw, 16/03/2022
Module A1


President
Aleksandra Kostrzewa



ANNEX TO THE CERTIFICATE

VALID ONLY WITH CERTIFICATE

No 1434-IVDD-035/2022

List of medical devices covered by the certificate:

Serial No.	Brand/Trademark	Product Name	REF. No.
1	ALLTEST	SARS-CoV-2 Antigen Rapid Test (Nasal Swab)	INCP-502H
2	Beright	SARS-CoV-2 Antigen Rapid Test (Nasal Swab)	INCP-502H
3	JusChek	SARS-CoV-2 Antigen Rapid Test (Nasal Swab)	INCP-502H
4	Lambra	SARS-CoV-2 Antigen Rapid Test (Nasal Swab)	INCP-502H
5	SCREEN CHECK TEST	SARS-CoV-2 Antigen Rapid Test (Nasal Swab)	INCP-502H
6	Rapid Response	SARS-CoV-2 Antigen Rapid Test (Nasal Swab)	INCP-502H
7	gruppo Si.Gi.	SARS-CoV-2 Antigen Rapid Test (Nasal Swab)	INCP-502H
8	AllChek	SARS-CoV-2 Antigen Rapid Test (Nasal Swab)	INCP-502H
9	NovaTec Immundiagnostica GmbH	GSD NovaGen SARS-CoV-2 Ag Rapid Test (Nasal Swab)	CVAG502H 05
10	Mila	Mila SARS-CoV-2 SZYBKIE TEST ANTYGENOWY (Wymaz z nosa)	INCP-502H
11	DNA DIAGNOSTIC	SARS-CoV-2 Antigen Rapid Test (Nasal Swab)	CV19AGH
12	BIOGNOST	CoviGnost AG Nasal HOME SARS-CoV-2 Antigen Rapid Test (Nasal Swab)	INCP-502H
13	INNOVA	SARS-CoV-2 Antigen Rapid Test (Nasal Swab)	INCP-502H
14	the one medical	SARS-CoV-2 Antigen Rapid Test (Nasal Swab)	INCP-502H



Issued under the Contract No. MD-34/2021
Application No: 048/2021
Certificate bears the authorized person signature.
Warsaw, 16/03/2022


President
Aleksandra Kostrzewa



**Add value.
Inspire trust.**

TÜV SÜD Product Service GmbH · Ridlerstr. 65 · 80339 Munich · Germany

Hangzhou AllTest Biotech Co., Ltd.
550#, Yin Hai street, Hangzhou Economic and Technologic Development
Area, 310018 Hangzhou
PEOPLE'S REPUBLIC OF CHINA

Your reference/letter of	Our reference/name	Tel. extension/Email	Fax extension	Date	Page
095123	GCN-SH251064A01; GCN-SH251064A02; GCN-SH251064A03; SH25106400_CLI	medical_devices@tuvsud.com		2025-05-06	1 of 8

**TÜV SÜD Product Service GmbH
Confirmation Letter
CLI 095123 0015 Rev. 00**

Reference: GCN-SH251064A01 | GCN-SH251064A02 | GCN-SH251064A03 | SH25106400_CLI

To whom it may concern,

Confirmation of the status of a formal application, written agreement, and appropriate surveillance in the framework of Regulation EU 2024/1860 amending Regulations (EU) 2017/746 (in the following referenced as IVDR) as regards the transitional provisions for certain in vitro diagnostic medical devices.

With this letter TÜV SÜD Product Service GmbH, designated under IVDR and identified by the number 0123 on NANDO, confirms that we have received a formal application in accordance with Section 4.3, first subparagraph of Annex VII of IVDR and has signed a written agreement in accordance with Section 4.3, second subparagraph of Annex VII of IVDR with the above stated manufacturer with the following Single Registration Number (SRN)

Single Registration Number: CN-MF-000010710

The devices covered by the formal application and the written agreement mentioned above are identified in the Tables below.

- Table 1 identifies the devices for which an IVDR application has been received, written agreement concluded and for which TÜV SÜD Product Service GmbH is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive.
- Table 2 identifies the devices for which an IVDR application has been received and a written agreement concluded, but TÜV SÜD Product Service GmbH has not yet taken the responsibility for appropriate surveillance of the corresponding devices under the applicable Directive or these devices did not require a Notified Body certificate under Directives.

Registered Office: Munich
Trade Register Munich HRB 85742
UniCredit Bank GmbH · BIC HYVEDEMMXXX
IBAN DE13 7002 0270 0048 8522 11
VAT ID No. DE129484267
Information pursuant to § 2 [1] DL-InfoV
(Germany) at tuvsud.com/imprint

Supervisory Board:
Holger Lindner (Chairman)
Board of Management:
Walter Reithmaier (CEO)
Patrick van Welij

TÜV SÜD Product Service GmbH
Zertifizierstelle für Medizinprodukte /
Certification Body for Medical Products
Ridlerstr. 65
80339 Munich
Germany

tuvsud.com/ps
Hotline: +49 89 50084-747





If devices covered by certificates issued under Directive 98/79/EC (IVDD) that expired after 26. May 2022 and before 09. July 2024, without having been withdrawn, this letter also confirms that

- the manufacturer signed the written agreement under IVDR by the date of IVDD certificate expiry; or
- provided evidence that a competent authority of a Member State had granted a derogation or exemption from the applicable conformity assessment procedure in accordance with Article 54(1) of IVDR or Article 92(1) of the IVDR respectively.

The transition timelines in accordance Article 110 (3) of IVDR that apply to the devices covered by this letter, subject to the manufacturer's continued compliance to the other conditions specified in Article 110 (3c) of IVDR, are shown below:

- 31. December 2027, for devices certified under IVDD
- 31. December 2027, for class D devices;
- 31. December 2028, for class C devices;
- 31. December 2029, for class B devices and for class A devices placed on the market in sterile condition

We reserve the right to invoice any issuance, copies, amendments and / or changes of the confirmation letter according to effort.

For confirmation letter validity see www.tuvsud.com/ps-cert?q=CLI 095123 0015

In case of inquiries please contact medical_devices@tuvsud.com.

The current revision of this Confirmation Letter is valid until **2025-09-26**.

On behalf of the Notified Body TÜV SÜD Product Service GmbH,
2025-05-06

TÜV SÜD Product Service GmbH
Medical and Health Services

TÜV SÜD Product Service GmbH
Medical and Health Services

Chenchuan Weng
[Chenchuan Weng \(May 6, 2025 15:20 GMT+8\)](#)

Mr. Chenchuan WENG
Conformity Assessment Responsible (CARE)

Michael Mauermeir
[Michael Mauermeir \(May 6, 2025 09:03 GMT+2\)](#)

Mr. Michael MAUERMEIR
Application Reviewer



Table 1: Devices covered by this letter and for which TÜV SÜD Product Service GmbH is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive:

Device name or Basic UDI-DI (under IVDR application)	IVDR Device classification (as proposed by the manufacturer and verified during application review)	If the IVDR device is a substitute device, identification of the corresponding IVDD device	IVDD Certificate Reference(s) of the devices under IVDR application, and the NB Identification
FSH Rapid Test Basic UDI-DI: 6970277510020PYK	Class B incl. ST/NPT	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
hCG Pregnancy Rapid Test Basic UDI-DI: 6970277510020QYM	Class B incl. ST/NPT	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
Digital hCG Pregnancy Test Basic UDI-DI: 6970277510020RYP	Class B incl. ST/NPT	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
hCG Pregnancy Rapid Test Basic UDI-DI: 6970277510020SYR	Class B incl. ST/NPT	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
hCG Pregnancy Enhanced Sensitivity Rapid Test Basic UDI-DI: 6970277510020TYT	Class B incl. ST/NPT	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
LH Ovulation Rapid Test Basic UDI-DI: 6970277510020UYV	Class B incl. ST/NPT	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
LH Ovulation Rapid Test Basic UDI-DI: 6970277510020VYX	Class B incl. ST/NPT	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
LH Ovulation Rapid Test Basic UDI-DI: 6970277510020WYZ	Class B incl. ST/NPT	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
LH Ovulation Rapid Test	Class B incl. ST/NPT	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00



Device name or Basic UDI-DI (under IVDR application)	IVDR Device classification (as proposed by the manufacturer and verified during application review)	If the IVDR device is a substitute device, identification of the corresponding IVDD device	IVDD Certificate Reference(s) of the devices under IVDR application, and the NB Identification
Basic UDI-DI: 6970277510020XZ3			NB# 0123
Chlamydia Rapid Test Basic UDI-DI: 6970277510020YZ5	Class C for professional use	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
CMV IgM Rapid Test Basic UDI-DI: 6970277510020ZZ7	Class C for professional use	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
H. pylori Antigen Rapid Test Basic UDI-DI: 6970277510020OYH	Class C incl. ST/NPT/CDx	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
Rubella IgM Rapid Test Basic UDI-DI: 6970277510021AXQ	Class C for professional use	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
Toxo IgG/IgM Rapid Test Basic UDI-DI: 6970277510021BXS	Class C for professional use	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
Toxo IgG/IgM Rapid Test Basic UDI-DI: 6970277510021CXU	Class C for professional use	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
ToRCH IgM Combo Rapid Test Basic UDI-DI: 6970277510021DXW	Class C for professional use	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
Vaginal pH Rapid Test Basic UDI-DI: 6970277510021EXY	Class C incl. ST/NPT/CDx	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
Ferritin Rapid Test Basic UDI-DI: 6970277510021FY2	Class C incl. ST/NPT/CDx	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
Sperm Concentration Rapid Test	Class B incl. ST/NPT	N/A	Certification as follows: V1 095123 0008 Rev. 04;



Device name or Basic UDI-DI (under IVDR application)	IVDR Device classification (as proposed by the manufacturer and verified during application review)	If the IVDR device is a substitute device, identification of the corresponding IVDD device	IVDD Certificate Reference(s) of the devices under IVDR application, and the NB Identification
Basic UDI-DI: 6970277510021GY4			VCQ 095123 0013 Rev. 00 NB# 0123
SP-10 Male Fertility Rapid Test Basic UDI-DI: 6970277510021HY6	Class B incl. ST/NPT	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
TSH Rapid Test Basic UDI-DI: 6970277510021IY8	Class C incl. ST/NPT/CDx	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
Vitamin D Rapid Test Basic UDI-DI: 6970277510021JYA	Class C incl. ST/NPT/CDx	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
FOB Rapid Test Basic UDI-DI: 6970277510020NYF	Class C incl. ST/NPT/CDx	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
PSA Rapid Test Basic UDI-DI: 6970277510021KYC	Class C for professional use	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
PSA Qualitative Rapid Test Basic UDI-DI: 6970277510021LYE	Class C for professional use	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
Urinary Tract Infections Test Basic UDI-DI: 6970277510019KXX	Class C incl. ST/NPT/CDx	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123
Urinary Tract Infections Test Basic UDI-DI: 6970277510019LYZ	Class C incl. ST/NPT/CDx	N/A	Certification as follows: V1 095123 0008 Rev. 04; VCQ 095123 0013 Rev. 00 NB# 0123

Legend: ST – self-testing; NPT – near-patient testing; CDx – companion diagnostics

Table 2: Devices covered by this letter and for which TÜV SÜD Product Service GmbH is NOT responsible for appropriate surveillance of the corresponding devices under the applicable Directive:

Device name or Basic UDI-DI (under IVDR application)	IVDR Device classification (as proposed by the manufacturer and verified during application review)	If the IVDR device is a substitute device, identification of the corresponding IVDD device	IVDD Certificate Reference(s) of the devices under IVDR application, and the NB Identification
HBsAg Rapid Test Basic UDI-DI: 6970277510017FYF	Class D incl. ST/NPT	N/A	Certification as follows: CeCert/102/W/E.1; CeCert/101/W/E.1; NB# 2934
HCV Rapid Test Basic UDI-DI: 6970277510017CY9	Class D incl. ST/NPT	N/A	Certification as follows: CeCert/107/W/E.1; CeCert/106/W/E.1; NB# 2934
HIV 1.2 Rapid Test Basic UDI-DI: 6970277510017BY7	Class D incl. ST/NPT	N/A	Certification as follows: CeCert/097/W/E.1; CeCert/096/W/E.1; NB# 2934
ABO and RhD Blood Grouping Rapid Test Basic UDI-DI: 6970277510021MYG	Class D incl. ST/NPT	N/A	Certification as follows: CeCert/089/W/E.1; CeCert/088/W/E.1; NB# 2934
SARS-COV-2 and Influenza A+B Antigen Combo Rapid Test Basic UDI-DI: 6970277510021NYJ	Class C incl. ST/NPT/CDx	N/A	Certification as follows: 1434-IVDD-217/2022; NB# 1434
SARS-CoV-2 Antigen Rapid Test Basic UDI-DI: 6970277510013OYM	Class C incl. ST/NPT/CDx	N/A	Certification as follows: 1434-IVDD-035/2022; NB# 1434
SARS-CoV-2/Influenza A+B/RSV/Adenovirus/M. pneumoniae Antigen Combo Rapid Test Basic UDI-DI: 6970277510021QYQ	Class B for professional use	N/A	N/A - Device did not require a Notified Body certificate under Directives
SARS-CoV-2 Antigen Rapid Test Basic UDI-DI: 6970277510021TYW	Class B for professional use	N/A	N/A - Device did not require a Notified Body certificate under Directives



Device name or Basic UDI-DI (under IVDR application)	IVDR Device classification (as proposed by the manufacturer and verified during application review)	If the IVDR device is a substitute device, identification of the corresponding IVDD device	IVDD Certificate Reference(s) of the devices under IVDR application, and the NB Identification
COVID-19 IgG/IgM Rapid Test Basic UDI-DI: 6970277510021SYU	Class B for professional use	N/A	N/A - Device did not require a Notified Body certificate under Directives

Confirmation Letter Version History

Date	TÜV SÜD Product Service GmbH internal reference traceable to each version of the letter	Action
2025-05-06	GCN-SH251064A01; GCN-SH251064A02; GCN-SH251064A03; SH25106400_CLI	Initial issue