

Q v Suchen: Alle Textspalten

Antigen-Tests zum direkten Erregernachweis des Coronavirus SARS-CoV-2

(i) Impressum

Liste der Antigen-Tests zum direkten Erregernachweis des Coronavirus SARS-CoV-2,

Los

die Gegenstand des Anspruchs nach § 1 Satz 1 gemäß "Dritte Verordnung zur Änderung der Verordnung zum Anspruch auf bestimmte Testungen für den Nachweis des Vorliegens einer Infektion mit dem Coronavirus SARS-CoV-2 (Coronavirus-Testverordnung – TestV)" sind.

Alle Daten gemäß Übermittlung des Herstellers, verbindlich sind ausschließlich die Angaben in den jeweiligen Gebrauchsinformationen.

Aktionen ~

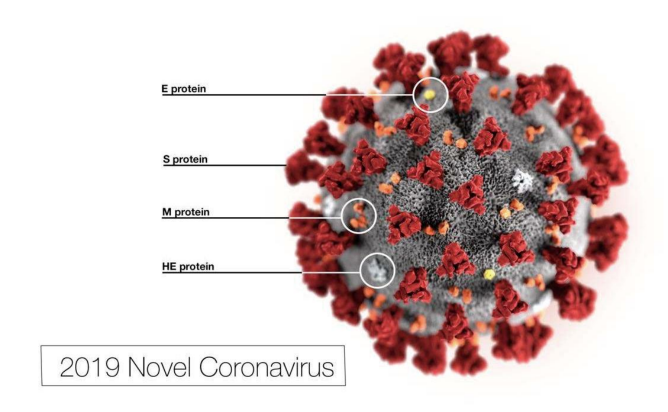
	Hersteller			Deutsche/r Vertreiber	Europäischer Bevollmächtigter						Sensitivität		Spezifität		
Test-ID	Name	Stadt	Land	Name	Name	Stadt	Land	Handelsname des Tests	Testort *	Artikelnummer	% ↓=	95%iges Vertrauens- intervall	%	95%iges Vertrauens- intervall	Gebrauchs- anweisung
AT174/20	JOYSBIO (Tianjin) Biotechnology Co., Ltd.	Tianjin	CN	ImuGeX GmbH	Lotus NL B.V.	The Hague	NL	SARS-COV-2 Antigen Rapid Test Kit (Colloidal Gold)	POC (ohne Gerät)	IGXJS-01	98,72	93,0 - 100,0	97,32	92,4 - 99,4	
AT046/20	JOYSBIO (Tianjin) Biotechnology CO., LTD.	Tianjin	CN		Lotus NL B.V.	The Hague	NL	SARS-CoV-2 Antigen Rapid Test Kit (Colloidal Gold) / Test Kit für neuartiges Coronavirus-Antigen (kolloidale Gold-Methode)	POC (ohne Gerät)	COV-AG-20 / G10313	98,72	93,0 - 100,0	97,32	92,4 - 99,4	S Link öffnen
AT184/20	JOYSBIO (Tianjin) Biotechnology Co., Ltd.	Tianjin	CN	New Mobility AG	Lotus NL B.V.	Den Haag	NL	SARS-CoV-2 Antigen Rapid Test Kit (Colloidal Gold) / Test Kit für neuartiges Coronavirus-Antigen (kolloidale Gold Methode)	POC (ohne Gerät)	COV-AG-20 / G10313	98,72	93,0 - 100,0	97,32	94,8 - 100,0	S Link öffnen
AT247/20	HANGZHOU CLONGENE BIOTECH CO., LTD.	Hangzhou	CN	Der Werbemarkt GmbH	Shanghai International Holding Corporation GmbH (Europe)	Hamburg	DE	CLUNGENE® COVID-19 Antigen Rapid Test	POC (ohne Gerät)		98,50	91,8 - 99,7	100,00	98,2 - 100	
AT181/20	Hangzhou Clongene Biotech Co., Ltd.	Hangzhou	CN	S2 Health GmbH	Shanghai International Holding Corporation GmbH (Europe)	Hamburg	DE	COVID-19 Antigen Rapid Test	POC (ohne Gerät)	04AGT19	98,50	91,8 - 99,7	100,00	98,2 - 100	
AT234/20	Hangzhou Clongene Biotech Co., Ltd.	Hangzhou	CN	Solenal GmbH	Shanghai International Holding Corp.GmbH (Europe)	Hamburg	DE	COVID-19 Antigen Rapid Test			98,50	91,8 - 99,7	100,00	98,2 - 100	S Link öffnen
AT099/20	Koch Biotechnology (Beijing) Co., Ltd.	Beijing	CN	enverque GmbH	Wellkang Ltd.	Dover	GB	COVID-19 Antigen Rapid Test Strip	POC (ohne Gerät)	NCV10	98,50	97,2 - 99,8	99,30	98,5 - 100	S Link öffnen
AT131/20	Hangzhou Clongene Biotech Co., Ltd.	Hangzhou	CN	Winlong GmbH	Shanghai International Holding Corp.GmbH (Europe)	Hamburg	DE	COVID-19 Antigen Rapid Test	POC (mit Gerät)		98,50	91,8 - 99,7	100,00	98,2 - 100	
AT183/20	Hangzhou Clongene Biotech Co., Ltd.	Yuang District Hangzhou	CN	Bauer -Kalrsruhe	Shanghai International Holding Corporation GmbH (Furone)	Hamburg	DE	COVID-19 Antigen Rapid Test	POC (ohne Gerät)		98,50	91,8 - 99,7	100,00	98,2 - 100	S Link öffnen

SARS-CoV-2 Antigen

PREFACE

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2, or 2019-nCoV) is an enveloped non-segmented positive-sense RNA virus. It is the cause of coronavirus disease 2019 (COVID-19), which is contagious in humans. SARS-CoV-2 has several structural proteins including spike (S), envelope (E), membrane (M) and nucleocapsid (N).

The antigen is generally detectable in upper respiratory samples during the acute phase of infection.

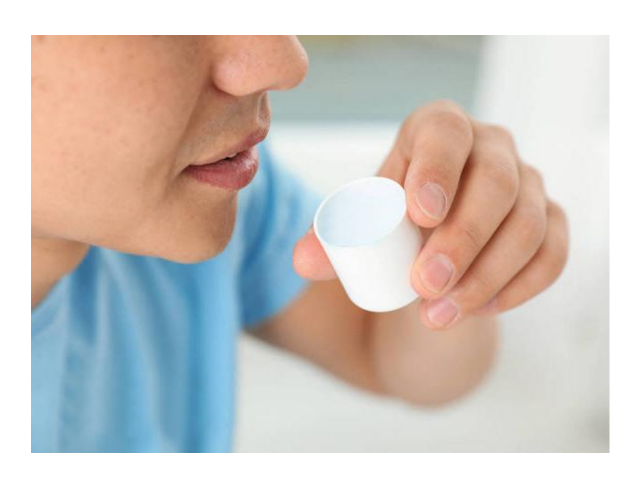


PRODUCT PHOTOS





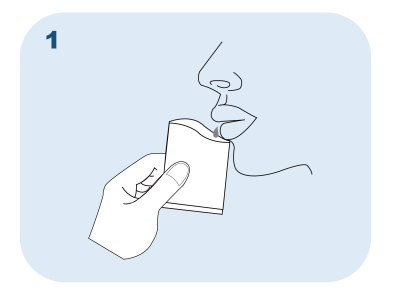




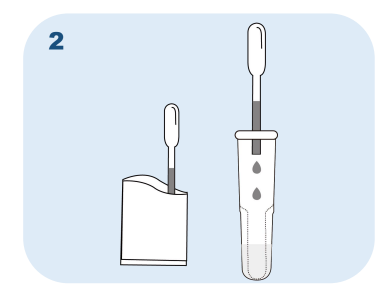
TEST PRINCIPLE

The Kit use immunocapture method, it is designed to detect the presence or absence of SARS-CoV-2 nucleocapsid proteins in respiratory samples from patients with signs and symptoms of infection who are suspected of COVID-19.

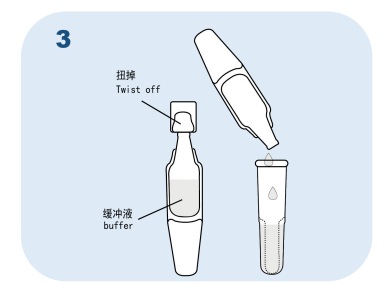
TEST METHOD



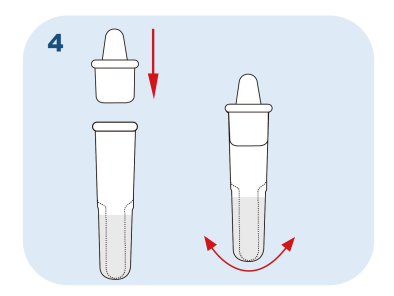
1. Before collecting oral fluid relax your cheeks and gently massage cheeks with fingers for 15-30 seconds, Gently spit oral fluid into the collection bag.



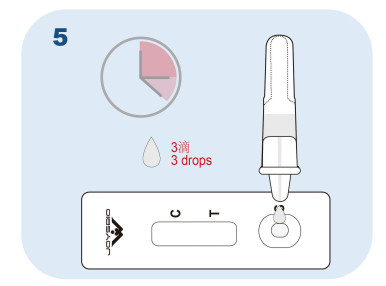
2. Hold the dropper vertically and draw oral fluid from collection bag and transfer 3 drops of oral fluid into the buffer bottle.



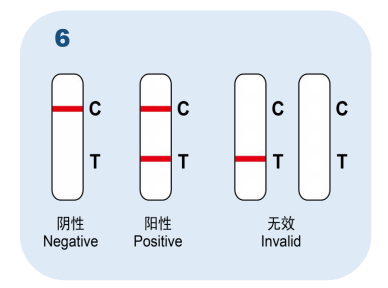
3. Twist off the top of the buffer bottle, slowly dispense all of the buffer into the extraction Tube.



4. Tighten the cap of the buffer bottle. Gently shake the buffer bottle for **10** seconds.



5. Tear off the foil pouch, take out the test strip/cassette and place the test kit on a clean and level surface. Gently squeeze the ridged body of the tube, dispensing three (3) drops of the processed specimen into the sample well. Read the test results between 15 and 20 minutes.



6.POSITIVE: Two lines appear. One colored line should be in the control line region (C), a colored line appears in test line (T) region. NEGATIVE: Only one colored control line appear. INVALID: Control line fails to appear.

CIINICAL EVALUATION REPORT

JOYSBIO SARS-CoV-2 Antigen Rapid Test Kit (Colloidal Gold) Clinical Evaluation Report

Consistency analysis of test results

There were 772 nasal swab specimens were collected to evaluate the clinical performance of the SARS-CoV-2 Antigen Rapid Test Kit Specimen Stability Study. The nasal swabs prospectively collected and enrolled from individual symptomatic patients (within 5 days of onset) who were suspected of COVID-19 and no duplicate samples were selected. Nasal swabs were collected following the dual nares method and handled as described in the package insert of the collection device.

A total of 154 samples were tested positive by SARS-CoV-2 Antigen Rapid Test Kit (Colloidal Gold). There were 2 samples in which the SARS-CoV-2 Antigen Rapid Test Kit ware positive and the Real-time fluorescent RT-PCR kit for detecting 2019-nCoV produced by BGI BIOTECHNOLOGY (WUHAN) ware negative, and 6 samples in which the SARS-CoV-2 Antigen Rapid Test Kit ware negative and the Real-time fluorescent RT-PCR kit for detecting 2019-nCoV produced by BGI BIOTECHNOLOGY (WUHAN) was positive.

There were 610 samples with negative test results in experimental reagent and 612 samples with negative test results in reference reagent. Hence, the sensitivity and specificity were 96.25% and 99.67% respectively.

The Real-Time Fluorescent RT-PCR Kit for Detecting SARS-CoV-2 produced by BGI BGI Genomics Co. Ltd was used as a comparator test. This is an FDA approved for EUA use product.

Overall Clinical Study Results

	PCR Co			
Reagent test results	positive	negative	Subtotal	
positive	154	2	156	
negative	6	610	616	
Subtotal	160	612	772	

Positive Percent Agreement (PPA)= 96.25% (95%CI:92.0%~98.6%)

Negative Percent Agreement (NPA)= 99.67% (95%CI:98.8%~100%)

Accuracy=98.96%

Kappa=0.97>0.5

Conclusion:

This clinical trial has performed a full analysis of the experimental reagents through methodological comparisons, and the results all meet the criteria for clinical evaluation. All the results showed that SARS-CoV-2 Antigen Rapid Test Kit (Colloidal Gold)meet the needs of clinical test.

SIGNIFICANCE

RESEARCH BACKGROUND

During the epidemic Situation, many countries have the following problems:

Existing detection methods cannot achieve large-scale rapid screening.

lack of technical expertise and inadequate laboratory capacity, Erroneous Operation can easily lead to missed inspections.

Can't afford high testing costs.





SIGNIFICANCE

According to the WHO, during the outbreak of SARS-CoV-2, in areas with confirmed SARS-CoV-2 community-wide transmission; confirmed outbreaks in closed or semi-closed communities; in high-risk groups; among contacts of confirmed cases; SARS-CoV-2 Antigen Rapid Test Kit (Colloidal Gold) as a tool to monitor disease incidence is a particularly effective detection method.

Globally, as of 3:59pm CEST, 17 August 2020, there have been 21,549,706 confirmed cases of COVID-19, including 767,158 deaths, reported to WHO.

ADVANTAGE

- 1.Easy to collect samples, simple operation, without professional equipment.
- 2. The test results are available in 15 minutes, and the test results are clearly visible.
- 3. Convenient transportation and low price, higher accuracy.
- 4. Suitable for large-scale rapid screening.



REGISTERED

REGISTERED







EU CE Certification

Emergency Use Authorization

WHO-Emergency Use Listing



> Retouradres Postbus 16114 2500 BC Den Haag

Lotus NL B.V. Tay de heer X Wei Koningin Julianaplein 10 2595 AA 's-Gravenhage

Datum: 18 augustus 2020

Betreft: aanmelding In-vitro diagnostica

Geachte heer Wei.

Op 13 augustus 2020 ontving ik uw notificatie krachtens artikel 4, eerste lid van Uw aanvraag het Nederlandse Besluit in-vitro diagnostica (BIVD) om onder de bedrijfsnaam 13 augustus 2020 JOYSBIO (Tianjin) Biotechnology Co., Ltd met Europees gemachtigde Lotus NL B.V. onderstaande producten als in-vitro diagnostica op de Europese markt te Correspondentie uitsluitend

De producten staan geregistreerd als in-vitro diagnostica onder nummer:

SARS-CoV-2 IgG/Neutralizing antibody Rapid Test Kit(Colloidal Gold) ,SARS-CoV-2 Antigen Rapid Test Kit (Colloidal Gold), Immunochromatography analyzer (geen merknaam) (NL-CA002-2020-53008)

Tuberculosis Antibody Test Kit (Colloidal Gold), Mycoplasma Pneumonia IgM Antibody Test Kit (Colloidal Gold), Treponema Pallidum Antibody Test Kit (Colloidal Gold), Morphine/Methamphetamine/Ketamine Test Kit (Colloidal Gold)

(geen merknaam) (NL-CA002-2020-53009)

Hiermee heeft u voldaan aan uw verplichting op grond van artikel 4, BIVD.

In alle verdere correspondentie betreffende bovenvermelde producten verzoek ik u deze nummers te vermelden. Aan deze nummers kunnen geen verdere rechten ontleend worden, ze dienen alleen om de notificatie administratief te vergemakkelijken.

De registratie van in-vitro diagnostica als medisch hulpmiddel op grond van de Classificatiecriteria (Bijlage II) bij Richtlijn 98/79/EG betreffende medische hulpmiddelen voor in-vitro diagnostiek is onderhevig aan mogelijke revisies van Europese regelgeving inzake de classificatie van medische hulpmiddelen en aan voortschrijdend wetenschappelijk inzicht (zie artikel artikel 10, eerste lid van Richtlijn 98/79/EG).

Farmatec

Bezoekadres: Riinstraat 50 2515 XP Den Haad T 070 340 6161

http://hulpmiddelen.farmatec.nl

Inlichtingen bij: M.P. Meiter - Michiels

medische_hulpmiddelen@ minvws.nl

Ons kenmerk:

Pagina 1 van 2

vermelding van de datum en het kenmerk van deze brief.

CE CERTIFICATE

Notificatie van in-vitro diagnostische medische hulpmiddelen impliceert dat de fabrikant, JOYSBIO (Tianjin) Biotechnology Co., Ltd de CE-conformiteitsmarkering heeft aangebracht op de desbetreffende producten alvorens deze in een EUlidstaat in de handel te brengen. Zodoende garandeert Lotus NL B.V. dat de invitro diagnostica voldoen aan de essentiële eisen zoals opgenomen in bijlage I bij Richtlijn 98/79/EG (en in het daarmee corresponderende onderdeel 1 bij het

Volledigheidshalve wijzen wij u erop dat een in-vitro diagnosticum moet voldoen aan de eisen uit het BIVD. Het BIVD is gebaseerd op Richtlijn voor in-vitro diagnostiek, 98/79/EG. Met name wijzen wij u op de Nederlandse-taaleis zoals deze in Nederland geldt, de eisen voor het ter beschikking houden van de technische documentatie en de plicht tot het hebben van een Post Marketing Surveillance- en vigilantiesysteem.

Tot slot merk ik op dat met uw notificatie - de administratieve notificatie als fabrikant - en deze brief geen sprake is van een oordeel over de status of kwalificatie van uw product; notificering betekent niet dat daadwerkelijk sprake is van een in-vitro diagnosticum in de zin van de onderhavige wet- en regelgeving. In voorkomende gevallen kan de Inspectie Gezondheidszorg en Jeugd (IGJ), belast met het toezicht op de naleving van het bij of krachtens de wet bepaalde, een standpunt innemen over de status van een product, waarbij het volgens vaste jurisprudentie uiteindelijk aan de nationale rechter is om te bepalen of een product onder de definitie van in-vitro diagnosticum valt.

De Minister voor Medische Zorg en Sport, namens deze,

Afdelingshoofd Farmatec

Dr. M.J. van de Velde

Pagina 2 van 2





Acknowledgment Letter

9/11/2020

Hongyan Li JOYSBIO (Tianjin) Biotechnology Co., Ltd. Tianjin Tianjin TEDA 300457 CHINA

Dear Hongyan Li:

The Center for Devices and Radiological Health (CDRH) of the Food and Drug Administration (FDA) has received your submission. This submission has been assigned the unique document control number below. All future correspondence regarding this submission should be identified prominently with the number assigned and should be submitted to the Document Control Center at the above letterhead address. Failure to do so may result in processing delays. If you believe the information identified below is incorrect, please contact the Office of Product Evaluation and Quality (OPEQ) submission support at (301) 796-5640 or OPEOSubmissionSupport@fda.hls.gov.

Submission Number: EUA202733

Received: 9/11/2020

Applicant: JOYSBIO (Tianjin) Biotechnology Co., Ltd. Device: SARS-CoV-2 Antigen Rapid Test Kit (Colloidal Gold)

We will notify you when the review of this document has been completed or if any additional information is required. For information about CDRH review regulations and policies, please refer to http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/default.htm.

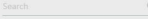
Sincerely yours,

Center for Devices and Radiological Health

U.S. Food & Drug Administration 10903 New Hampshire Avenue Silver Spring, MD 20993 www.fda.gov

Has entered the FIND recommended list











COVID-19

WHO WE ARE Y

WHAT WE DO Y

NEWSROOM

CALLS FOR PARTNERS

- Hunan Yonghe-Sun Biotechnology Co., Ltd SAKS-COV-2 specific antibody test kit (Immunochromatography) (RUO) Contact
- InDevR Inc. COVID Serology Kit: Multiplexed Immunoassay (RUO) Contact
- Innovita Biological Technology Co. Ltd 2019-nCoV Antibody Test (Colloidal Gold) (China NMPA EUA Australia TGA Brazil ANVISA Singapore HSA CE-IVD)
- InTec Products, Inc. Rapid SARS-CoV-2 Antibody Test (CE-IVD) Contact 1 Contact 2
- InTec Products, Inc. Rapid SARS-CoV-2 Antibody (IgM/IgG) (CE-IVD) Contact 1 Contact 2
- Jetta Labs LLP OZO Diamond SARS-CoV2 (COVID-19) IgG/IgM Test (Latex Method) (CE-IVD) Contact
- Jetta Labs LLP OZO India SARS-CoV2 (COVID-19) IgG/IgM Test (Colloidal Gold Method) (CE-IVD) Contact
- Jiangsu Bioperfectus Technologies Co. Ltd PerfectPOC Novel Corona Virus (SARS-CoV-2) IgM/IgG Rapid Test Kit (CE-IVD) Contact
- Jiangsu Bioperfectus Technologies Co. Ltd PerfectPOC Novel Corona Virus (SARS-CoV-2) Aq Rapid Test Kit (CE-IVD) Contact
- Jiangsu Superbio Biomedical Technology (Nanjing) Co., Ltd SARS-CoV-2 (COVID-19) IgM/IgG Antibody Fast Detection Kit (Colloidal Gold) (US FDA EUA CE-IVD)
- JinHuan Medical Instrument Co., Ltd (COVID-19) IgM/IgG Antibody Fast Detection Kit (Colloidal Gold) (CE-IVD) Contact
- Joinstar Biomedical Technology Co., Ltd SARS-CoV-2 IgM/IgG Antibody Test (Colloidal Gold) (CE-IVD) Contact
- JOYSBIO (Tianjin) Biotechnology Co., Ltd COVID-19 IgG/IgM Rapid Test Kit (Colloidal Gold) (CE-IVD) Contact
- JOYSBIO (Tianjin) Biotechnology Co., Ltd COVID-19 (SARS-CoV-2) Antigen Rapid Test Kit (Colloidal Gold) (CE-IVD) Contact
- JOYSBIO (Tianjin) Biotechnology Co., Ltd COVID-19 Neutralizing Antibody Test Kit (Lateral Flow Rapid Test) (CE-IVD) Contact
- Kephera Diagnostics KDx Rapid SARS-CoV-2 Antigen Test (In development) Contact
- Kephera Diagnostics KDx COVID-19 IgG/IgM Rapid Detection Test Kit (In development) Contact
- Koch Biotechnology (Beijing) Co., Ltd SARS-CoV-2 Antigen Lateral Flow Assay (MHRA UK) Contact
- KRISHGEN BioSystems Human Anti-SARS-CoV-2 (Covid-19) IgG/IgM Rapid Test (CE-IVD) Contact
- KRISHGEN BioSystems Human Anti-SARS-CoV-2 (Covid-19) IgM Rapid Test (RUO) Contact
- L&H Biotech Limited COVID-19 Antiqen Rapid Test (In development) Contact
- Labnovation Technologies Inc. COVID-19 (SARS-CoV-2) IqM/IqG Antibody Test Kit (CE-IVD) Contact 1 Contact 2
- Labtest Diagnostica SA Anti COVID-19 IgG/IgM Rapid Test (Brazil ANVISA) Contact
- Leadgene Biomedical, Inc. Leadgene® SARS/SARS-CoV-2 Antigen Rapid Test Kit (In development) Contact
- Leadgene Biomedical, Inc. Leadgene ® SARS/SARS-CoV-2 IqG/IqM Rapid Test Kit (In development) Contact
- <u>Lifeassay Diagnostics Pty Ltd</u> Test-it COVID-19 IgM/IgG Lateral Flow Assay (In development) <u>Contact</u>
- . LifeSensors, Inc. COVID-19 IgG ELISA Detection Kit (RUO) Contact
- Liming Bio-Products Co., Ltd COVID-19 IgG/IgM Combo Rapid Test Device (CE-IVD) Contact
- LOMINA AG Fast COVID19 IgM/IgG Antibody Detection Kit (Colloidal Gold) (CE-IVD) Contact
- Luminostics, Inc. CLIP-COVID19 (smartphone-read out high sensivity antigen detection test) (In development) Contact

Search Website

https://www.finddx.org/cov id-19/pipeline/

The Emergency Use Listing



SARS-CoV-2 Rapid Antigen Tests: progress of the active applications in the emergency use listing assessment pipeline

Product name	Product code(s)	Manufacturer name	Dossier review	QMS Desk Assessment	
ESPLINE SARS-CoV-2	231906	Fujirebio, Inc	R		
BIOEASY Diagnostic kit for SARS-CoV-2 Ag (Fluorescence Immunochromatographic Assay)	YRLF04401025, YRLF04401050 and YRLF04401100	Shenzhen Bioeasy Biotechnology Co., Ltd	awaiting submission	awaiting submission	
LumiraDx SARS-CoV-2 Ag Test	L0160001nnxxx	LumiraDx UK Ltd	awaiting submission	awaiting submission	
SARS-CoV-2 Rapid Antigen Test	9327592190	Roche Diagnostics GmbH			
SARS-CoV-2 Antigen Rapid Test Kit (Colloidal Gold)	G10313	JOYSBIO (Tianjin) Biotechnology CO., LTD			

Progress of the active applications in the emergency use listing assessment pipeline.

JOYSBIO (Tianjin) Biotechnology Co., Ltd.

COMPANY PROFILE

JOYSBIO (Tianjin) Biotechnology Co.,

Ltd. is a Chinese R&D-based biotechnology company that develops, manufactures, and supplies high-quality medical in-vitro diagnostic (IVD) rapid test kits as well as revolutionary customized solution kits to all parts of the world. Founded by a team of professionals with many years of combined technical,marketing/sales, operational and manufacturing expertise in this industry, we offer high quality but cost-effective rapid test kit.

