# Iumasis

COVID-19 lgG/lgM Test

One Step COVID-19 lgG/laM Test

11( Please read the instructions carefully before use!

#### INTENDED USE ]

Humasis COVID-19 IgG/IgM test is one step in vitro diagnostic test based on an immunochromatographic assay. It is designed for qualitative detection of Immunoglobulin G and Immunoglobulin M antibody of Novel Coronavirus (COVID-19) in human blood.

#### SUMMARY AND EXPLANATION ]

Coronavirus is a group of viruses that belongs to the Family Coronaviridae; a type of RNA virus of 27~32kb commonly found in birds and mammals including human. Coronavirus is divided into four genera: alpha, beta, gamma and delta. The virus causes illness ranging from the common cold to more severe diseases such as Middle East Respiratory Syndrome (MERS-CoV) and Severe Acute Respiratory Syndrome (SARS-CoV).

Coronavirus disease 2019 (COVID-19) is a new strain caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2). The disease originated from Wuhan city of China in December 2019. The World Health Organization (WHO) publicly named this virus 'COVID-19' and declared it a pandemic and a Public Health Emergency of International Concern. The infection typically spreads from one person to another via direct contact or respiratory droplets from cough or sneeze. Latent period from exposure to onset of symptoms is between one to fourteen days (four to seven days on average). Common symptoms and signs of infection include fever, cough, shortness of breath and breathing difficulties. In severe cases, infections can cause pneumonia, severe acute respiratory syndrome, kidney failure and even death. Due to the wide variety of symptoms, it is difficult to differentiate COVID-19 from other existing respiratory viruses or bacteria.

Diagnosing COVID-19 through isolating the virus or detecting specific genes from the collected respiratory droplet specimens is a challenge in terms of time and accessibility as it requires long hours, well-equipped laboratory and advanced technology which are often not available to many public. Therefore the need for point-of-care rapid test kit that can conduct preliminary screening of COVID-19 which requires less time and cost is rising.

# [ PRINCIPLE OF THE TEST ]

"Humasis COVID-19 IgG/IgM Test" is a rapid immunochromatographic assay test which detects IgG and IgM antibody to COVID-19 in human blood. A nitrocellulose membrane strip in the device contains two test lines (G and M line) and a control line (C). G line is pre-coated with mouse anti-human lgG for detection of lgG anti-COVID-19, and M line is pre-coated with mouse anti-human IgM for detection of IgM anti-COVID-19. C(Control line) is coated with goat anti-mouse IgG.

When sample is added to sample pad, it moves through the conjugate pad, where recombinant antigen-colloidal gold particle will react with the IgG and IgM antibodies specific to COVID-19 in the sample, forming an immunocomplex. The complex moves along the membrane by capillary action and makes contact with the immobilized antibody coated in the test region. Colored line in the test region indicates a positive result for coronavirus. The absence of colored line in the test region suggests a negative result. The complex continues to move to the control region and will react with immobilized reagents that capture colored conjugate regardless of test specimen composition. The resulting visible colored line in the control region confirms that the assay is functioning correctly and its result is valid.

# CONTENTS ]

- 1. Humasis COVID-19 IgG/IgM Test device
- 2. Assay diluent
- 3. Instruction for use
- Optional: Capillary tube (10uL)

## MATERIAL COMPOSISTION 1

#### 1 test device contains :

Mouse anti-human IgM monoclonal antibody	0.44±0.11 μg
Mouse anti-human IgG monoclonal antibody	0.44±0.11 μg
2019-nCOV n recombinant protein	0.08±0.02 μg
Goat anti-mouse IoG	$0.08 \pm 0.02 \ \mu q$

## STORAGE AND SHELF-LIFE ]

Store the test device packaged in a sealed foil pouch at 2 to 30°C (36~86°F) 2. Shelf-life : 6 months from manufacturing date

#### [SPECIMEN COLLECTION AND PREPARATION ]

- 1. The device can be performed using whole blood, plasma, serum and finger puncture whole blood
- 2. Whole blood : Collect specimen in collection tube with anticoagulant such as EDTA, heparin or sodium citrate. Perform test immediately after collection, or can be stored at 2~8°C up to 24 hours before the test.
- 3. Plasma : Collect specimen in collection tube with anticoagulant and centrifuge the sample. The sample can be stored at 2~8°C up to 3 days, and freeze the sample for longer storage.
- 4. Serum : Collect specimen in collection tube without anticoagulant and place it in room temperature for 30 minutes before the centrifuge. Serum sample can be stored at 2~8°C up to 3 days, and freeze the sample for longer storage.
- 5. Finger puncture whole blood : Collect the sample with lancet on clean fingertip. Avoid squeezing the fingertip as it will dilute the blood sample and use immediately after collection.

## [ TEST PROCEDURE ]

- 1. If the collected specimens were stored in refrigerated condition, leave the samples in room temperature for 15 to 30 minutes before the test. Avoid unsealing the device if the device temperature is lower than the room temperature.
- 2. Open the sealed pouch and place the device on a clean, dry and level surface.
- 3. Release 10uL of whole blood, plasma or serum into the sample well.
- Then add 2~3 drops (70~100uL) of sample diluent immediately.
- 4. Read the result at 15 minutes. Do not read result after 15 minutes.



## [INTERPRETATION OF RESULT ]

#### 1. Negative

If no colored line appears in the test region (G, M) and a colored line is present on the control region (C), then the result is negative.



#### 2. Positive

0

In addition to the presence of colored line in control region(C)





IqG positive : if there is colored line in G but no development in M, then the result is IgG positive.



M), then the result is positive.

then the result is IgM positive.

IgM positive : if there is no colored line in G but visible colored line in M,

#### 3. Invalid

If there is no colored line in the control region (C), the result is invalid.



# [ PRECAUTIONS AND LIMITATIONS ]

- For in vitro diagnostic use only
  Do not use the test device beyond the expiration date.
- 3. Keep sealed until usage, and once opened use immediately.
- 4. Do not use the test device if the pouch is damaged or the device is seriously broken.
- 5. Do not re-use the device.
- 6. Handle all specimens safely as potentially infectious.
- 7. This test is intended for initial screening of coronavirus infection by detecting antibody to COVID-19, but should not be used as a sole criterion for the determination of coronavirus infection. Other methods and clinical information (signs and symptoms) should be used and considered for diagnose.

## [ REFERENCES ]

- Korean Centers for Disease Control http://ncov.mohw.go.kr/ -
- FIND https://www.finddx.org/covid-19/
- CDC https://www.cdc.gov/
- Development and Clinical Application of a Rapid IgM-IgG Combined Antibody Test for SARS-CoV-2 Infection Diagnosis. Z Li,. Journal of Medical Virology

<b>IVD</b> : For <i>in vitro</i> diagnostic use	<b>LOT</b> : Lot number	<b>REF</b> : Catalogue number
<b>I</b> : Consult instructions for use	$^{30^{\circ}C}$ : Store at 2~30°C	() : Do not reuse
<b>ECREP</b> : Authorized Representative	: Manufactured by	: Use by / Expiry date
CE: This product fulfills the requirements for Directive 98/79/EC on <i>in vitro</i> diagnostic medical devices		

CE



Humasis Co., Ltd. Rm. 114, 502, 504, 604, 604-1, B03-1, B03-2, 88, Jeonpa-ro, Dongan-gu, Anyang-si, Gyeonggi-do, 14042, Republic of Korea TEL:+82-31-478-8597, FAX:+82-31-478-8586 Email:question@humasis.com www.humasis.com

**ECREP** MT Promedt Consulting GmbH Altenhofstr. 80 D46386 St. Ingbert / Germany TEL: +49 6894 - 58 10 20 FAX: +49 6894 - 58 10 21 Email: info@mt-procons.com www.mt-procons.com