

## Instructions for Use (IFU)

# SARS-CoV-2/COVID19 IgM/IgG Antibody Fast Detection Kit (Colloidal Gold)-PROFI (Not intended for internal use)



IVD

SÚKL / RZPRO / EUDAMED  
# 061583-0001 # 061583-0002

GMDN

#64756

### [Product name]

SARS-CoV-2/COVID19 IgM/IgG Antibody fast detection kit (Colloidal gold) - PROF I

### [Intended use]

20 or 5 doses/tests (each dose contains a testing tape with colloidal gold and cards showing the batch code, date of manufacture, and expiry date).

### [Intended use]

IVD test for IgM and IgG antibodies in blood, i.e. for proving the Coronavirus SARS-CoV-2 infection/ COVID-19 disease. Edition for Health Care Professionals

### [Test principle]

This test is based on fluorescent immunochromatography. The testing kit contains 1) an antigen labelled by colloidal gold and a complex of control antibodies; 2) nitrocellulose membranes with two testing lines (M- and G-line) and one qualitative control line (C-line). The required amount of the sample is dripped into a hole on the test strip. Capillary effects help the sample run down the nitrocellulose membrane inside the test strip. If the tested sample contains SARS-CoV-2/COVID-19 IgM/IgG antibodies, they will be bound to the COVID-19-antigen labelled by colloidal gold and the monoclonal IgM or IgG antibodies will appear on the nitrocellulose membrane as violet-red M- or G-lines. The test will show the sample positivity to IgM or IgG antibodies, thus proving the COVID-19 infection.

### [Content of presentation]

- ① Plastic testing strip
- ② 2 plastic droppers
- ③ Vial with reagent
- ④ Lancet ("vial") for blood sampling
- ⑤ Disinfecting pad
- ⑥ Instructions for use

### [Storage conditions]

① Store under the temperature of from 4 to 30 °C. Protect from light. The expiry date and the batch number are shown on the primary pack of the test.

② Use immediately after opening the primary pack. The test starts degrading due to humidity after the primary pack opening. The test cannot be used 60 minutes after opening; the results will not be valid.

### [Usable biological material]

The test has been developed and tested for blood samples testing by the general public.

[Additional information for health professionals: Peripheral blood, serum or plasma samples obtained with clinically used anticoagulants (EDTA, heparin, sodium citrate) can be used for the test.]

## [Instructions for use]

Open the box, take the primary pack out and let it warm/cool to the room temperature (24 °C). At first, read the whole instructions for use and use the kit immediately after you open the primary pack, however, within maximum 60 minutes after you open the aluminium primary pack



Lancet ("vial") for blood sample taking (contains a needle)



Plastic droppers



Disinfecting pad



Vial with reagent



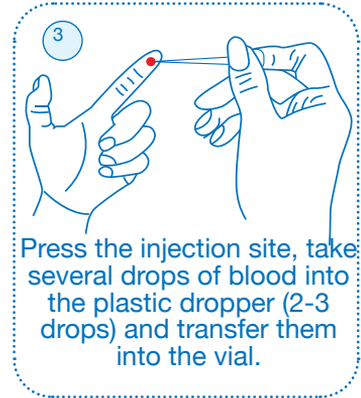
Plastic testing strip



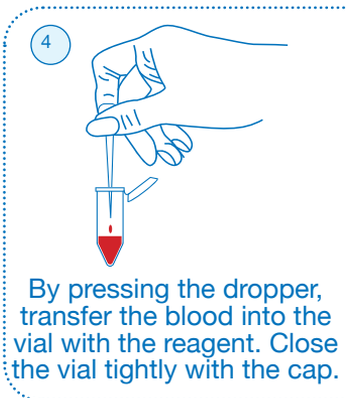
1  
Clean the blood sampling site with a disinfecting swab and let it air-dry.



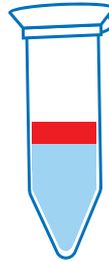
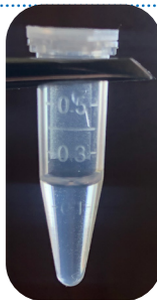
2  
Remove the lancet cap and press the narrower lancet end at the disinfected site to release the needle.



3  
Press the injection site, take several drops of blood into the plastic dropper (2-3 drops) and transfer them into the vial.



4  
By pressing the dropper, transfer the blood into the vial with the reagent. Close the vial tightly with the cap.



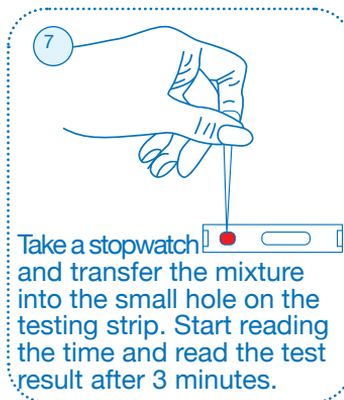
Fill in the vial by 2-3 drops!  
Otherwise the test may be invalid!



5  
Check whether the vial is closed tightly and mix the mixture well by shaking (for approx. 30 seconds)



6  
Use a clean dropper and draw the mixed mixture in it; use 2-3 drops exactly



7  
Take a stopwatch and transfer the mixture into the small hole on the testing strip. Start reading the time and read the test result after 3 minutes.



8  
Dry the excess blood, if any, and disinfect the site again. Pack the test remains and dispose them in a prescribed way.

**ATTENTION!** Do not read the test result after more than 10 minutes! The test result can change due to humidity and oxidation! The results are invalid after 10 minutes!

The results are valid only from minute 3 to minute 10 after the mixture dripping on the plastic testing strip. Let the strip remain in a horizontal position at rest. Do not move or tilt it during the measurement! Use immediately after opening the primary pack!

**HINT:** Take a photo of the strip and then read the result.

## [Test results reading]

### 1. Negative result:

If the QCC (Quality Control – C) control line is visible and the M- and G-lines are not, the sample is negative because no antibodies were detected. However, continue your health condition monitoring. Re-testing 24 hours later is recommended. Do not forget that the amount of antibodies produced in case of a disease is individual and antibodies production is manifested usually by increased temperature.

#### ATTENTION!

The test detects the presence of antibodies but its precision depends on the amount of antibodies which is individual. Even if the test is negative, please monitor your health condition. If increased temperature and flu-like symptoms persist, repeat the test 24-48 hours later. If respiratory or any other serious complaints occur, or if your health condition does not improve, consult your physician. If you have symptoms and you stayed in infected environment or were in contact with an infected individual, always consult your condition and actions with your physician.

### 2. Positive test result:

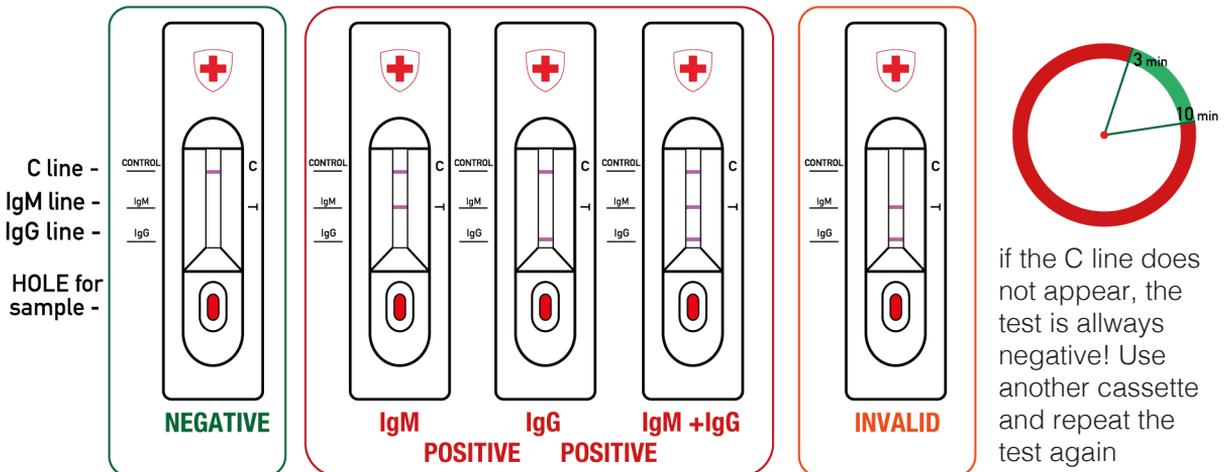
2.1 If the control C-line appears together with the M-line, IgM antibody was detected and the sample is positive.

2.2 If the control C-line appears together with the G-line, IgG antibody was detected and the sample is positive.

2.3 If the control C-line appears together with both the G- and M-lines, both IgG and IgM antibodies were detected and the sample is positive. Stay calm in all three cases and read the instructions on page 4.

### 3. Invalid result:

If no C-line appears, the test is invalid and must be repeated! It can be due to time delay at testing, reagent contamination, test impairment by temperature etc.



Dispose of infectious / contaminated material according to standard procedures.

### [Cross-reactivity]

Tests for IgG / IgM antibodies against the SARS-CoV-2 virus (whole blood / serum / plasma) were analyzed to detect antibodies in positive samples of influenza virus type A, B, RSV, adenovirus, HBsAg, syphilis, H pylori, HIV and HCV. The results did not show any sign of cross reactivity.

### [Disruptive substances]

The following compounds were tested with the SARS-CoV-2 IgG / IgM antibody test (whole blood / serum / plasma) and no interfering effects were found. Triglycerides: 50 mg / dl, ascorbic acid: 20 mg / dl, hemoglobin 1000 mg / dl, bilirubin: 60 mg / dl, total cholesterol: 6 mmol / l

The results did not show cross reactivity.

### [specificity / sensitivity]

96,4% / 99,7%

