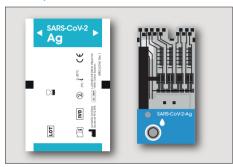
Quick Reference Instructions

Warning and Precautions:

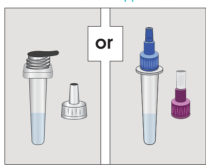
All kit components can be discarded as Biohazard waste according to local guidelines. Refer to the product safety data sheet for risk and safety phrases and disposal information. The product safety data sheet is available at lumiradx.com. Exercise the normal precautions required for handling all laboratory reagents. Proper laboratory safety techniques should be followed at all times when working with SARS-CoV-2 patient samples. Patient swabs, used Test Strips and used extraction buffer vials may be potentially infectious. Proper handling and disposal methods should be established by the laboratory in accordance with local, state and federal regulations. Reagents encapsulated within the Test Strip are present in extremely small amounts and where any component is of animal origin, the source is certified as free from infectious or contagious material – however, should any reagent become exposed it should be treated as potentially infectious.

LumiraDx SARS-CoV-2 Ag Test Kit Components

Test Strip



Extraction Vial and Dropper Lids



The LumiraDx SARS-CoV-2 Ag Test is rapid microfluidic immunofluorescence assay for use with the LumiraDx Platform intended for the qualitative detection of the nucleocapsid protein antigen to SARS-CoV-2 directly from nasal swab samples collected from individuals suspected of, or at increased risk of, COVID-19 by their healthcare provider within the first twelve days of symptom onset.

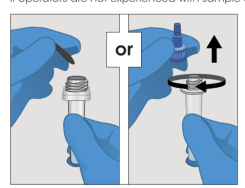
Study the LumiraDx Platform User Manual and LumiraDx SARS-CoV-2 Ag Test Strip Product Insert thoroughly before using these Quick Reference Instructions or performing a test. This is not a complete package insert.

Operate the LumiraDx Platform at room temperature between 15°C and 30°C (59°F and 86°F) and 10% - 90% relative humidity. The extracted sample must be used within 5 hours of preparation when stored at room temperature. Extracted nasal samples may be frozen at -80°C and used up to 5 days after freezing. Samples and extraction buffer must be at room temperature before testing. Check expiration date on outer test kit carton and each individual test package before using. **Do not use any test beyond its expiration date**. Refer to the LumiraDx SARS-CoV-2 Ag Test Strip Product Insert for sample collection instructions. Refer to the LumiraDx SARS-CoV-2 Ag Test Strip Product Insert for Sample Collection, Warning and Precautions, and Limitations.

Preparing the sample

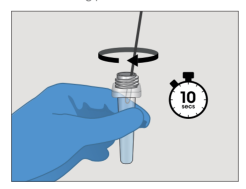
Collect a patient swab sample before following steps 1 - 4 of Running the Test.

Sample Collection and Handling: Proper sample collection and handling of nasal swabs is required to ensure accurate results (refer to product insert). Additional training or guidance is recommended if operators are not experienced with sample collection and handling procedures.



Remove seal

Remove the seal <u>or blue screw cap</u> from top of **Extraction Vial** containing the **Extraction Buffer**.



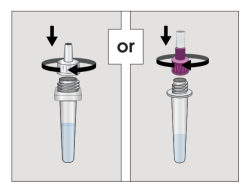
Soak Swab

Place and soak the **Patient Swab** in the **Extraction Buffer** for 10 seconds then stir well by rotating the swab against the side of the vial 5 times.



Squeeze Swab

Remove the **Patient Swab** while squeezing the **Extraction Vial** to remove the liquid from the swab Discard the swab in biologard waste



Attach Dropper Lid

Firmly attach the <u>clear or purple</u> **Dropper Lid** to the top of the **Extraction Vial**. The extracted sample must be used (see Step 5 and 6 below) within 5h of preparation when stored at room temperature.

Cleaning and Disinfecting

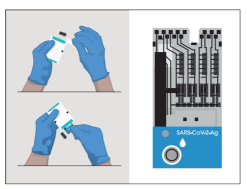
Wipe the external surfaces of the LumiraDx Instrument with a soft, slightly damp cloth when it appears visibly dirty. Disinfect the Instrument after each patient test or if contamination is suspected using LumiraDx approved materials. Details of LumiraDx approved disinfectant materials can be found at lumiradx.com. Use the material until the surface of the Instrument is visibly wet. Allow the surface to remain wet for 1 minute and let air dry. Avoid USB ports and power inlet.

Do not spray or pour solution directly onto the Instrument. Do not put any objects or cleaning materials into the Test Strip slot.

Running the Test



1. Select Patient Test from the Instrument Home Screen and enter patient details using the Keyboard or Barcode Scanner. See section 10 of the Platform User Manual for instructions on using the Barcode Scanner.

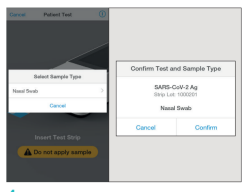


2. Remove the **Test Strip** from its pouch and hold by gripping only the blue portion. **Do not bend** the **Test Strip** or touch any part other than the blue portion.

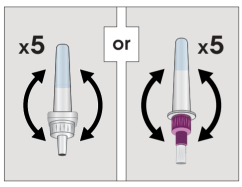


Patient Test

3. When prompted, open the Instrument door and gently insert the Test Strip as far as it will go. The thick black alignment rib on the Test Strip should be on the left and line up with the black line on the Instrument. Do not apply the sample until prompted. Install the Lot Calibration file if using a new Test Strip Lot for the first time. See section 2.8 of the Platform User Manual.



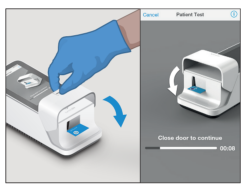
4. Select the appropriate sample type and confirm the test type.



5. Gently invert the **Extraction Vial** five times just before applying the sample to the **Test Strip**.



6. Apply **one whole drop** of the sample onto the **Test Strip Sample Application Area** when prompted by the **Instrument**.



7. Close the door when prompted to continue the test.



8. Results are displayed within 12 minutes of applying the sample. The left-hand image here shows a positive result for SARS-CoV-2 Ag and the right-hand image shows a negative result for SARS-CoV-2 Ag. Tap *Finish* to complete testing or tap *Comment* to leave a comment or to reject the Test, then follow prompts to return to the *Home Screen*. All test results must be read using the Lumiradx Instrument.

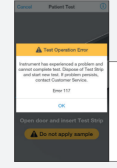
INTERPRETATION OF RESULTS

Positive Test Results:

SARS-CoV-2 antigen present; does not rule out coinfection with other pathogens.

Negative Test Results:

Negative results are presumptive for patients with symptom onset beyond 12 days. Negative test results do not preclude infection and should not be used as the sole basis for treatment or other patient management decisions, including infection control decisions, particularly in the presence of clinical signs and symptoms consistent with COVID-19, or in those who have been in contact with the virus. It is recommended that these results be confirmed by a molecular testing method, if necessary for patient management.



Example of an error screen:
If the On Board Control (OBC)
fails, an error message will be
shown and no test result will be
returned. Follow the on screen
instructions to dispose of the
Test Strip and start a new test. If
the problem persists, contact
Customer Services.

Invalid Results:

If an issue occurs, a message will be displayed on the Instrument touch-screen. Alert messages include useful information and are highlighted by an orange banner. Error messages also include a $\stackrel{\bullet}{\mathbf{A}}$ symbol. All messages will contain a description of the Instrument status or error and an instruction. Error messages contain an identifying code that may be used for further troubleshooting purposes.

Quality Controls

To complete Quality Control assessment of the LumiraDx Instrument and SARS-CoV-2 Ag Test Strips, you must use the LumiraDx SARS-CoV-2 Ag Quality Control Pack which are available separately. If the LumiraDx Antigen Quality Controls do not perform as expected, do not report patient results. Retest using a new Test Strip – if problems persist contact LumiraDx Customer Services.

Customer Service

If the LumiraDx SARS-CoV-2 Ag Test or the LumiraDx Instrument do not perform as expected, contact LumiraDx Customer Services via lumiradx.com or customerservices@lumiradx.com



Manufacturer Information

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