For use under the Emergency Use Authorization (EUA) only For in vitro diagnostic use

INTENDED USE

The Sampinute[™] COVID-19 Antigen MIA is intended for the qualitative detection of spike proteins of SARS-CoV-2 in nasopharyngeal swab (NPS) specimens in conjunction with the Sampinute[™] Analyzer. The test cartridge, with the analyzer, incorporates a technique called magnetic force-assisted electrochemical sandwich immunoassayⁱ. The technique entails the use of magnetic nanoparticles (MNPs) and electrochemical sensors conjugated with monoclonal antibodies specific for the receptor binding domains (RBD) of SARS-CoV-2 spike proteins to facilitate the electrochemical sandwich immunoassay. Testing is limited to laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C. §263a, that meet the requirements to perform high complexity tests.

SARS-CoV-2 spike proteins are generally detectable in NPS specimens during the acute phase of infection. Hence, results reveal the presence or absence of SARS-CoV-2 viral antigens in the NPS specimens. Positive results indicate the presence of viral antigens, but clinical correlations with patient history and other diagnostic information are necessary to determine infection status.

Positive results do not rule out bacterial infection or co-infection with other viruses. The agent detected may not be the definite cause of disease. Laboratories within the United States and its territories are required to report all positive results to the appropriate public health authorities.

Negative results should be treated as presumptive and confirmed with a molecular assay, if patient management is necessary. Negative results do not rule out the possibility of COVID-19 infection and should not be used as the sole basis for treatment or patient management decisions, including infection-control decisions. Negative results should be considered in the context of a patient's recent exposures, history and clinical signs and symptoms consistent with COVID-19.

The Sampinute[™] COVID-19 Antigen MIA is intended for use with Sampinute[™] Analyzer by medical professionals or trained operators who are proficient in the use of the device. Both the Sampinute[™] Analyzer and the Sampinute[™] COVID-19 Antigen MIA are only for use under the Food and Drug Administration's Emergency Use Authorization.

SUMMARY AND EXPLANATION

SARS-CoV-2, which causes the disease COVID-19, was first discovered in Wuhan, Hubei Province, China in December 2019, where it was first thought to have spread from bats. However, the coronavirus is also thought to have infected other animals as intermediary hosts.^{II} The virus spreads primarily through small droplets produced from coughing, sneezing, and talking.^{III} The median incubation time is estimated to be 5.1 days with symptoms expected to be present within 12 days of infection.^{IV}

PRINCIPLE OF THE TEST

The Sampinute[™] COVID-19 Antigen MIA employs magnetic force-assisted electrochemical sandwich immunoassay that is used with Sampinute[™] Analyzer to detect spike proteins from SARS-CoV-2.

The recommended period of incubation until the test is equivalent to that of diagnostic PCR testing for SARS-CoV-2. (It can be found that the virus has a median incubation time of approximately 5.1 days. Symptoms usually manifest after being infected for 12 days.) The patient sample is collected using a NPS and is either directly placed into the Reagent Tube supplied in the kit or a universal viral transport system. In the case of the Reagent Tube, the sample should be processed for analysis within one hour. As for the universal viral transport system, room temperature-equilibrated samples can be directly processed for analysis. If the transport system is frozen, the sample should be fully thawed and equilibrated to room temperature.

The sample is dispensed into the sample inlet of a Sampinute[™] COVID-19 Antigen MIA test cartridge.

The sample flows along the microfluidic channel, and if SARS-CoV-2 spike proteins are present in the sample, they form complexes with anti-SARS-CoV-2 spike protein antibodies conjugated to MNPs. These complexes eventually encounter and bind onto the working electrode of the electrochemical sensor, that is coated with anti-SARS-CoV-2 spike protein antibodies. *Via* magnetic actuation, the antigen-antibody reactions are actively controlled. This ensures that the MNPs and the antigens are thoroughly mixed to form immuno-complexes on the electrode. The unbound MNPs are removed *via* magnetic field.

Subsequently to the magnetic actuation step, the device proceeds to the introduction of a detection buffer followed by the electrochemical measurement step, wherein a voltage is applied to induce electrochemical oxidation and reduction of gold on the MNPs, resulting in the electric current.

The test results reveal the presence (positive) or absence (negative) of SARS-CoV-2 spike protein antigens. The Sampinute[™] Analyzer will display the test results (positive, negative) on the screen.

REAGENTS AND MATERIALS SUPPLIED

Principal Ingredients

Each test cartridge of the Sampinute[™] COVID-19 Antigen MIA contains the main reactants and components shown below:

- MNPs conjugated with monoclonal antibodies specific for receptor binding domain (RBD) of spike protein from SARS-CoV-2.
- Electrodes coated with monoclonal antibodies specific for RBD of spike protein from SARS-CoV-2.

Product Package Components

25 test cartridges of Sampinute[™] COVID-19 Antigen MIA are provided in the package, which can be used to analyze up to 25 samples, including the control solutions provided in the package.

The package contains the following:

- Sampinute[™] COVID-19 Antigen MIA (25): test cartridges with monoclonal anti-SARS-CoV-2 antibodies, MNPs, and electrochemical sensors.
- Reagent Tubes (25): solutions for collecting specimens.
- Sterile nasal swabs (25): flexible swabs for collecting specimens.
- Negative control solution (1): salt solution with less than 0.1% sodium azide. Upon measurement, the results are expected to show negative, as no spike proteins are present in the control solution. One valid result of the analysis of the negative control solution should be obtained per package.
- Positive control solution (1): salt solution with non-infectious SARS-CoV-2 antigen and less than 0.1% sodium azide. Upon measurement, the results are expected to show positive, as SARS-CoV-2 spike proteins are present in the control solution. One valid result of the analysis of the positive control solution should be obtained per package.
- Package insert (1)

MATERIALS NOT SUPPLIED

- Sampinute[™] Analyzer
- Adapter
- Power cord
- Barcode scanner
- Printer
- BD Universal Viral Transport (UVT), Copan Universal Transport Media (UTM), or Viral Transport Media (VTM), the preparation of which follows the CDC protocol
- A pipette and disposable tips to transfer the specimen (or a glass capillary)
- A vortex shaker (or a mixer)

WARNINGS AND PRECAUTIONS

- Only use Sampinute[™] COVID-19 Antigen MIA test cartridges with the Sampinute[™] Analyzer.
- Do not use expired Sampinute[™] COVID-19 Antigen MIA test cartridges.
- Do not use Sampinute[™] COVID-19 Antigen MIA test cartridges that are damaged or broken.
- The recommended measurement temperature range is 15-30°C (59-86°F).
- Only use the Sampinute[™] COVID-19 Antigen MIA test cartridges after a full system check according to the Sampinute[™] Analyzer user manual.
- Wear disposable protective gloves when handling the Sampinute[™] COVID-19 Antigen MIA test cartridges and human specimens.
- If transport of the samples is required, the following transport media have been tested and shown not to interfere with the performance of the test.
 - BD[™] Universal Viral Transport System
 - Copan[®] UTM-RT[®] System
 - CDC VTM according to SOP#: DSR-052-02
- Gently press the Sampinute[™] COVID-19 Antigen MIA test cartridge down onto the tray of the Sampinute[™] Analyzer gently until it will go no further.
- The product is for single use only. Do not re-use.
- Place the test cartridge on the tray with the labelled side facing up.
- Take a precise amount of sample for each test and insert it into the sample inlet at once.
- Discard the Sampinute[™] COVID-19 Antigen MIA test cartridge after use as required by internal quality control procedures and in accordance with local, state and federal regulations or accreditation requirements.
- Do not swallow or damage the Sampinute[™] COVID-19 Antigen MIA test cartridge.
- The product is for *in vitro* diagnostics.
- The product is indicated for use in clinical laboratories.
- Follow safety procedures set by your institution for working with chemicals and handling biological specimens.
- This test has not been FDA-cleared or -approved. The test has been validated, but FDA's independent review of this validation is pending.
- This test has been validated only for the detection of proteins from SARS-CoV-2, not for any other viruses or pathogens.

KIT STORAGE AND STABILITY

Keep the cartridge packaged in the provided aluminium pouch. Keep the product refrigerated (2-8°C, 36-46°F). **Upon preparation, the test cartridge must be placed at room temperature**

(15-30°C, 59-86°F) at least 30 minutes before use. Use the cartridge immediately after opening the aluminum pouch. Do not expose the product to direct sunlight.

QUALITY CONTROL

Before the actual sample test, the Sampinute[™] Analyzer and the Sampinute[™] COVID-19 Antigen MIA test cartridges must go through a system check, as well as an external quality control test using positive and negative sample control solutions.

Sampinute[™] Analyzer Test Internal Controls

The screen on the device will show whether the sample analyzed is either negative, positive or associated with any error(s).

All test procedures (e.g. sample injection, antigen enrichment, buffer insertion) are monitored *via* internal sensors, where errors or improper executions are communicated through automated error messages.

No.	Error Notifications	Descriptions and Solutions
1	'Please input the admin password'	Admin password is not inputted.
		Displayed when touching the 'DONE' button without entering the admin password.
		Solution: Input the admin password and then touch the 'DONE' button.
		Hospital code is not inputted.
2	'Please input the hospital code.'	Displayed when touching the 'DONE' button without entering the hospital code.
		Solution: If a hospital code is required, contact the administrator. Input the hospital code and then touch the 'DONE' button.
3	'Unregistered hospital code. Please contact administrator.'	Unregistered hospital code
		Displayed when unregistered hospital code is entered.

Table 1:Error notifications, their descriptions and solutions

No.	Error Notifications	Descriptions and Solutions
		Solution: If a hospital code is required, contact the administrator. Input registered hospital code. If the issue persists, contact the administrator.
		No response from server
4	'Failed to connect to network. Please try again.'	Displayed when device registration fails due to lost network connection or other errors which make server communication unavailable.
		Solution: Either try to connect again at a later time or connect to another network.
		Wi-Fi communication error
5	'Unknown error. Please try again.'	The alert pops up when Wi-Fi communication is failed.
		Solution: Try again. If the issue persists, then contact the administrator.
	'Please try again after connecting to a network.'	No Wi-Fi connection
6		Displayed when device requests server communication for Wi-Fi connection.
		Solution: Connect to Wi-Fi.
	'Low battery. 15% of battery is remaining. The remaining battery must be more than 15% to run the analyzer. Please connect the analyzer to the power source.'	Low battery
7		Displayed when touching the low-battery icon on the home screen.
		Solution: Connect to a power adapter.
8	'The system check was failed or has an error. Please check the results again. Contact the system administrator.'	System Check is not performed. Displayed when system check is not performed.
		[Setting screen → System Check → Run System Check] Solution: Perform system check.
		QC not performed.
9	'The QC was failed or has an error. Please check the results again.'	Displayed when touching 'QC!' icon on the home screen in the case of when the QC Test has not been

No.	Error Notifications	Descriptions and Solutions
		performed.
		Solution: Perform QC test.
		Administrator account logout
10	'The admin account automatically logged out.'	Displayed when administrator account logs out before power off or reboot.
	'Please log in first.'	Attempted access in a logged off state
11		Displayed when the setting menu requires administrator account login.
		Solution: Log into an administrator account.
		Master password has not been entered.
12	'Please input the master password.'	Displayed when the 'DONE' button is touched without master password input.
		Solution: Enter the master password.
		Password input error
13	'Incorrect password. Please input the master password again.'	Displayed when the wrong password has been entered.
		Solution: Enter the correct password.
	'The fan is activated due to increased internal temperature of the analyzer.'	Fan activated
14		Displayed when the fan is activated by Sampinute™ Analyzer's temperature sensor operation.
		Wrong operator ID
15	'Invalid ID. Please check the ID again.'	Displayed when wrong operator ID is entered.
		Solution: Enter the correct ID.
		Operator ID has not been entered.
16	'Please input the operator ID.'	Displayed when touching DONE without entering the operator ID.

No.	Error Notifications	Descriptions and Solutions
		Solution: Enter the operator ID
		Patient code has not been entered
17		Fatient code has not been entered.
	'Please input the patient code.'	Displayed when touching DONE without entering the patient code.
		Solution: Enter the patient code.
		Tray ejection failure
18	'Please insert test cartridge in the tray.'	Displayed when a cartridge is not inserted in the tray within the specified time period.
		Solution: Insert the cartridge immediately upon request.
19		Cartridge is not removed.
	'Please remove the cartridge.'	Displayed when cartridge is not removed upon request.
		Solution: Remove the cartridge.
20		Current password mismatch.
	'Incorrect password. Please input the password correctly.'	Displayed when you do not enter currently set password.
		Solution: Enter the currently set password.
		New password mismatch
21	'The new password and re- entered password do not match. Please input the passwords again.'	Displayed on the Change Administrator Password screen when newly entered password does not match with the one typed in the new password confirm box.
		Solution: Ensure both passwords entered are the same.
22		New administrator password has not been entered.
	'Please input the new admin password.'	Displayed when a new administrator password has not been entered in the password setting screen.
		Solution: Enter new administrator password.

No.	Error Notifications	Descriptions and Solutions
23		Bluetooth disconnection
	'Please disconnect the device first and then try connecting the device.'	Displayed when a new Bluetooth printer connection is requested while the analyzer is connected to a Bluetooth printer.
		Solution: Connect to the appropriate Bluetooth printer.
24		USB disconnection
	'Please remove the USB port first and then try connecting the device.'	Displayed when a Bluetooth printer connection is requested while the analyzer is connected to a USB mobile printer.
		Solution: Either disconnect the Bluetooth or the USB printer.
25		Inappropriate temperature
	The temperature is not within operational range. (Current temperature: 38°C / 100.4°F)	Displayed when touching the temperature icon on the home screen which pops up when temperature is out of range for testing.
		Solution: Re-locate device in an environment of operational range (15-30°C / 59-86°F).

Table 2: Error pop-ups, their descriptions and solutions

Error pop-ups	Descriptions and Solutions
	Inappropriate temperature
Out Of Temperature Range	Case1.
Current temperature 38°C (100.4°F)	Pops up when the 'RUN TEST' button is touched on the home screen, or the 'RUN QC TEST' button is touched.
The temperature is not within operational range. For accurate readings, please try again at room temperature.	Case2. Pops up when the temperature is out of test range.
ок	Solution: Move the device to the recommended

Error pop-ups	Descriptions and Solutions
	operating temperature (15-30°C, 59-86°F) and wait until device temperature decreases to below 30°C (86°F).
	Low battery
EXAMPLE TO BATTERY LOW! Low battery. 15% of battery is remaining. The remaining battery must be more than 15% to run the analyzer. Please connect the analyzer to the power source.	Case1. Pops up when the 'RUN TEST' button is touched on the home screen, or the 'RUN QC TEST' button is touched. Case2. Pops up when the battery level is 15% or lower and the user touched Next to proceed to the next level of QC test.
	Solution: Connect the analyzer to a power source.
Image: Construct of the state of the st	QC test has not been performed. Pops up when the 'RUN TEST' button on the home screen is touched under the condition that QC TEST has not been performed. Solution: Run the QC TEST.
QC Test Error! The QC test was failed or has an error. Please test again or contact the system administrator.	QC test error Pops up when the 'RUN TEST' button is touched on the home screen under the condition that QC lock settings are turned 'ON', and/or QC TEST result is 'Fail'.
ОК	Solution: Conduct QC test again. If the QC test is a fail, then contact the system administrator.

Error pop-ups	Descriptions and Solutions
EXAMPLE Run System Check First Please perform the system check. The system check should be performed prior to sample tests. OK	A system check has not been performed. Pops up when the 'RUN TEST' button is touched on the home screen under the condition that the system check has not been performed. Solution: Perform system check.
	System Check Error
System Check Error! The system check was failed or has an error. Please test again or contact the system administrator	Pops up when the 'RUN TEST' button is touched on the home screen under the condition that the system check result is 'Fail'.
ОК	Solution: Conduct system check again. If the system check is a fail, then contact the system administrator.
Contact System Administrator	Contact the system administrator.
If you forgot your admin password, contact the system administrator. - e-mail: support@bbbtech.com	Pops up upon selecting 'Forgot password?'
- Tel: +82-070-4407-8808	Solution: Contact the system administrator.
Error!	Invalid cartridge QR code
Invalid QR code. Please scan the test cartridge QR code again.	Pops up when an invalid QR code is scanned.
ОК	Solution: Scan a valid QR code.

Error pop-ups	Descriptions and Solutions
Error! The test cartridge has been expired. Please use a new test cartridge. OK	Expired cartridge QR code Pops up when the QR code of the expired cartridge is scanned. Solution: Scan a new cartridge of valid expiry date.
Error! The wrong type of cartridge scanned. Please check the type of cartridge and scan the cartridge QR code again.	The QR code of the check cartridge is scanned. Pops up when the check cartridge's QR code is scanned instead of the test cartridge. Solution: Scan the QR code of a test cartridge.
Error!(ER1) Please contact the system administrator. OK	Tray ejection failure Pops up within 10 seconds after tray ejection fails. Solution: Contact the system administrator.
Error! The test cartridge has already been used. Please use a different test cartridge. OK	Pops up when a used cartridge is inserted. Solution: Remove used cartridge and insert an unused cartridge.

Error pop-ups	Descriptions and Solutions
	Insufficient sample
L Insufficient Sample Please try again with enough volume (at least 50 μL). If failure persists for the third time, try again with a new cartridge. ΟΚ	Pops up when a sample test fails due to errors associated with timeout, insufficient test sample injection, etc. Solution: Insert at least 50 μL of the sample again into the inlet. If failure persists for the third time, insert a new cartridge.
Cover Open Printer cover is open. Please close the cover and try again. OK	Pops up when the printer cover is open. Solution: Close the printer cover.
Paper Empty Please load paper and try again.	Pops up when there is no paper left in the printer. Solution: Re-fill printer with paper.
ОК	
Printer Disconnected Please check the Connection and try again.	Printer disconnection Pops up when the printer is not connected and the user touches the 'Print' button.
ОК	Solution: Ensure that the printer is connected.

Error pop-ups	Descriptions and Solutions
ER101 There was an error proceeding the analysis. Please retry the test or contact the system administrator.	Analysis failure Pops up when analysis is failed. Solution: Either try the analysis again or contact the administrator.
Line for the set of t	Insufficient solution Pops up when a solution test fails due to either timeout, insufficient solution injection, or solution reading failure. Solution: Insert at least 50 μL of the control solution again into the inlet. If failure persists for the third time, insert a new cartridge.
Error! The wrong type of cartridge scanned. Please scan check cartridge QR code.	QR code of the test cartridge is scanned. Pops up when the QR code of the test cartridge is scanned during the System Check process. Solution: Scan the QR code of a check cartridge.
Version Update Failed New version update failed. Please try again. OK	New version installation failure Pops up upon new version installation failure during software update. Solution: Re-try version update.

Error pop-ups	Descriptions and Solutions
	USB Device Connection Error
USB Device Recognition Error Connect a USB device and try again in 5 seconds. OK	The alert pops up when USB is removed or disconnected during software update. Solution: Connect the USB device and try again in 5 seconds.
Update Failed Firmware update failed. Would you like to try again? TRY AGAIN	Pops up when the firmware update failed. Solution: Try updating the firmware again.
	Repetitive update failure
Update Failed Firmware update failed. Please contact system administrator.	Pops up when firmware updates failed more than 3 times.
ок	Solution: Contact the system administrator.
Battery Low!	Low battery during software update
Battery too low to allow update. Please connect the charger and update when battery level is more than 15%.	Pops up upon touching 'Update' when battery level is 15 % or lower.
ОК	Solution: Connect the device to a charger.

Sampinute[™] Analyzer System Check Procedure

System Check: The purpose of this test is to verify the proper operation of the Sampinute[™] Analyzer.

- 1. Touch the 'Run System Check' button
- 2. Scan the QR code of the check cartridge
 - a. Scan QR code on the check cartridge with the barcode scanner.
 - b. After scanning the QR code, the screen will display "Insert Check Cartridge."

Note:

- The barcode scanner has to be purchased separately.
- If you scan the QR code of the test cartridge, the test will not proceed.
- If the QR code recognition continues to fail, please contact the administrator.
- 3. Insertion of the check cartridge tray
 - a. Insert the check cartridge to the tray according to instructions on the screen.
 - b. Insert the check cartridge in the correct position.
 - c. Touch the **"OK"** button.





- 4. Start system check
 - a. The system check takes about 1 minute.
- 5. Confirm the system check results
 - a. The test results are displayed upon completion.
 - b. After checking the results, remove the used cartridge from the tray.
 - c. Touch the **"DONE"** (DONE) button to move to the home screen.

Note:

Pass : Indicates that the device is working properly and a " ν " (\bigcirc) mark is displayed on the screen.

Fail : An error message and an "X" (X) mark is displayed on the screen.

Try out the test again or contact the administrator.

If the test fails under [Settings \rightarrow Lock Settings \rightarrow QC fail \rightarrow ON], a test cannot be run.

System Check → System Check Results

The System Check Results are displayed in order from oldest to newest.



Sampinute[™] Analyzer Test External Quality Control Test

The purpose of the external quality control test is to ensure that the test kit properly differentiates the positive and negative samples before the test of the patient specimen.

It is recommended that controls are run once:

- for each different test cartridge lot,
- for each new operator,
- as considered by internal quality control procedures and in accordance with local, state and federal regulations or accreditation requirements.

If either additional quality control solutions are needed or the procedure does not perform as expected, contact Celltrion Customer Services or BBB Customer Services.

- BBB Inc. (via website: <u>www.bbbtech.com</u>)
- Celltrion USA, Inc. (via email: Sampinute@celltrion.com, or via phone: (201) 988-4615)
- 1. Touch the **"Run QC Test"** button.
- 2. Scan the operator ID.
 - a. Scan the operator ID with the barcode scanner or manually enter the operator ID using the keypad. Touch the **"OK"** (OK) button only if entered manually.

Note: If you do not have a barcode scanner, touch **"Input operator ID manually"** to enter the code using the keypad.



- 3. Scan the test cartridge QR code
 - a. Scan the QR code on the cartridge pouch with the barcode scanner.
 - b. Upon scanning the QR code, the test level, cartridge type and LOT will display on the screen.

Note: The barcode scanner has to be purchased separately. If the QR code recognition fails, scan a new cartridge pouch. Please check if the cartridge type is correct.

- 4. Insert the test cartridge
 - a. Ensure the correct operator ID is entered.
 - b. Check that you scanned the QR code of the cartridge to be inserted.
 - c. Double check whether you are using the test cartridge for the correct target analyte.
 - d. Before inserting the cartridge, ensure that you are not using a used one.
 - e. When the error message, **"Please contact the system administrator"** is displayed after inserting the cartridge, stop the process and contact the administrator to resolve the issue.



- 5. Inject the positive control solution
 - a. Swirl or flick the bottom of the tube for 15 seconds to ensure thorough mixing.
 - b. Add 2-3 drops of the positive control solution into the sample inlet of the test cartridge within two minutes.
 - c. Touch the **"OK"** (OK) button.
- 6. Discard the used cartridge.
 - a. Refer to the images below to eject the cartridge from the tray. Discard the Sampinute[™] COVID-19 Antigen MIA test cartridge after use as required by internal quality control procedures and in accordance with local, state and federal regulations or accreditation requirements.



b. Touch the **"OK" (OK)** button.



- c. When touching the **"OK"** (**OK**) button, the screen will proceed to the **"QR Code Scan"** screen.
- 7. Test cartridge QR code scan
 - a. Scan the QR code on the cartridge pouch with the barcode scanner.
 - b. After scanning the QR code, the test level, cartridge type and LOT will display on the screen.

Note: The barcode scanner has to be purchased separately. If the QR code recognition fails, scan a new cartridge pouch. Please check if the test marker is correct.

- 8. Insert the test cartridge
 - a. Ensure that the correct operator ID is entered before inserting the cartridge.
 - b. Check that you scanned the QR code of the cartridge to be inserted.
 - c. Double check whether you are using the test cartridge for the correct target analyte.
 - d. Before inserting the cartridge, ensure that you are not using a used one.
 - e. When the error message, "Please contact the system administrator" is displayed on the screen after inserting the cartridge, stop the process and contact the administrator to resolve the issue
- 9. Negative control solution injection
 - a. Swirl or flick the bottom of the tube for 15 seconds to ensure thorough mixing.
 - b. Add 2-3 drops of the negative control solution into the sample inlet within two minutes.



- c. Touch the **"OK" (OK)** button.
- 10. Check the result of the QC Test
 - a. When the test is completed, the result of the QC test is displayed on the screen.
 - b. Check the result and remove the cartridge from the tray.
 - c. Touch the "DONE" (DONE) button → Return to the home screen.

Note: If the test results show "Pass", then successful operation of the test cartridge is indicated. If test results show "Fail", then the accuracy of the test cannot be verified, and the test needs to be conducted again or the administrator should be contacted.

- 11. Discard the used cartridge.
 - a. Refer to the images below to eject the cartridge from the tray.
 - b. Discard the Sampinute[™] COVID-19 Antigen MIA test cartridge after use as required by internal quality control procedures and in accordance with local, state and federal regulations or accreditation requirements.



c. Touch the **"DONE"** (DONE) button.



SAMPLE COLLECTION AND HANDLING SAMPLE COLLECTION

Nasopharyngeal Swab Specimen



Insert the sterile nasal swab through the nostril parallel to the palate (not upwards) until resistance is encountered or the distance is equivalent to that from the ear to the nostril of the patient, indicating contact with the nasopharynx. The swab should reach depth equal to the distance from nostrils to the outer opening of the ear. Gently rub and roll the swab. Leave the swab in place for several seconds to absorb secretions. Slowly remove the swab while rotating it.

A Swab Test Procedure in Reagent Solution Tube

Place swabs immediately and directly into sterile tubes containing the reagent solution. To collect a sample:

- 1. Aseptically take off and discard the cap from the tube.
- Insert the swab into the tube. Swirl the swab in the solution for 15 seconds. Plunge the swab in vertical motion for at least another 15 seconds in the solution. Ensure that the solution does not splash out of the tube when swirling and plunging.
- 3. Remove and discard the swab while pressing against the sides of the tube to ensure maximum amount of liquid has been squeezed from the swab.
- 4. Assemble and press the tip firmly onto the reagent solution tube containing the specimen. Mix thoroughly by either flicking the bottom of the tube or swirling.





B Swab Test Procedure in Viral Transport Media

Place the swab immediately and directly into the Viral Transport System or Universal Transport Media System following their package instructions for use inserted.

- 1. Check that the transport media tube cap is closed.
- 2. If frozen specimens were prepared, melt the frozen specimens completely before the test.
- 3. Mix thoroughly by either flicking the bottom of the tube or swirling. If a vortex mixer can be used, vortex for at least 1 minute the collection tube to ensure thorough mixing.

Warning:

- **1**. Do not use tips, tubes or caps taken from other products.
- 2. If frozen specimens were prepared, melt the frozen specimens completely before the test.

SAMPLE TRANSPORT AND STORAGE

The reagent solutions should be stored at room temperature (15-30°C, 59-86°F) upon testing. It is recommended that the specimens upon collection are processed and analyzed as soon as possible. The specimen would only be viable for processing for up to one hour.

For UVT, UTM, VTM and transport media prepared according to the CDC protocol, store specimens at room temperature (15-30°C, 59-86°F) for up to 4 hours and at 2-8°C (36-46°F) for up to 72 hours after collection. If a delay in testing or shipping is expected, store specimens at - 70°C (-94°F) or below.

TEST PROCEDURE

Run the test at a temperature range of 15-30°C (59-86°F). If the temperature is out of the range, a test cannot be run. When the battery level is 15% or lower, you cannot start the test. Fully charge the battery before running the test.

Expiration date: Check the expiration date on each individual test package or outer box before using. *Do not use any tests past the expiration date on the label.*

1. Touch the "**RUN TEST"** button to start the test

a. Put on a clean pair of gloves.

- b. Touch the **"RUN TEST"** button.
- 2. Scan the operator ID

a. Scan the operator ID with the barcode scanner or manually enter the operator ID using the keypad. Touch the **"OK"** (OK) button only if entered manually.

3. Scan the patient code

a. Scan the patient code with the barcode scanner or manually enter the code using the keypad. Touch the **"OK"** (OK) button only if entered manually.

b. The operator ID and the patient code are displayed on the screen. The screen subsequently proceeds to the screen for QR code scan.

4. Scan the QR code of the test cartridge

a. Scan the QR code on the cartridge pouch with the barcode scanner.

b. After scanning the QR code, the cartridge type and LOT code will be displayed on the screen, and the predetermined "cut-off" value of the cartridge (shown in page 27) is also

recognized, which is unchangeable and varies depending on the lot. (The cut-off value for each lot is pre-determined *via* selection and analysis of a few sample cartridges of each lot. The cut-off value defines the threshold signal for distinguishing negative and positive results.)

Subsequently, the screen proceeds to cartridge insertion.

(The barcode scanner has to be purchased separately. If the QR code recognition fails, scan a new cartridge pouch. After scanning, please ensure that the cartridge type displayed on screen is Sampinute[™] COVID-19 Antigen MIA.)

5. Test cartridge insertion

a. Take out the Sampinute[™] COVID-19 Antigen MIA test cartridge from the aluminum pouch.

b. Insert the test cartridge to the tray following instructions on the screen.

c. When the test cartridge is inserted in the correct position, the screen automatically proceeds to the sample injection stage.



Note:

- Make sure to check whether the correct operator ID and patient codes are entered before scanning the QR code.
- Ensure that the correct test cartridge is being used for the target analyte.
- Before inserting the cartridge, ensure that you are not using an already used one.
- When the error message, "Please contact the system administrator" is displayed after inserting the cartridge, stop the process and contact the administrator to resolve the issue.
- 6. Inject sample to the test cartridge using the reagent solution included in the package

a. Prepare the sample according to "Sample Collection – *Nasopharyngeal Swab Specimen*" on page 22.

b. Ensure that the specimen is mixed thoroughly by either flicking the bottom of the tube or swirling.

c. Add 2-3 drops of the sample into the inlet of the Sampinute[™] COVID-19 Antigen MIA test cartridge by inverting the tube and holding it vertically approximately an inch above the sample inlet, squeezing the bottom of the tube gently. (Injected sample needs to fill up the inlet of the cartridge, and excess volume can be used for further testing if required.)

d. Touch the **"OK"** (OK) button.

7. Inject sample to the test cartridge using the Transport Media.

a. Prepare the sample according to "Sample Collection – Nasopharyngeal Swab Specimen" on page 23.

b. Set the volume of the micropipette at 50μ L and attach disposable tip.

c. Remove cap on the collection tube.

f. Withdraw the sample in the collection tube with the micropipette. Check that the pipette tip does not contain bubbles.

- c. Draw 50 μL of the specimen and insert it into the inlet of the cartridge.
- d. Touch the **"OK"** (OK) button.
- 8. Process Analysis and Test Results
 - a. The progress of the analysis is shown as a percentage.
 - 1) The screen displays the image below to direct the usage of the reagent solution.



2) The screen displays the image below, in the case of using transport media.



b. When the analysis is completed, the result of the measurement is displayed on the screen with the patient code, operator ID, cartridge type and the LOT, and the tray at the bottom of the device is ejected.

- c. Remove the used test cartridge from the tray following **STEP 8**.
- d. Touch the **"DONE"** (**DONE**) button.

Note:

• Do not turn off or unplug the power adaptor while a test is in progress.

9. Discard the used cartridge.

- Refer to the images below to eject the cartridge from the tray.
- Discard the Sampinute[™] COVID-19 Antigen MIA test cartridge with any components used after use as required by internal quality control procedures and in accordance with local, state and federal regulations or accreditation requirements.



10. Test Results

a. Results of past tests can be viewed by touching the **"Test Results"** button on the home screen.

b. All test results are displayed in the order from the newest to oldest scan.

c. You can select specific results to view details.

d. If a printer is connected, you can print the results by clicking the print icon.



Warning: If refrigerated cartridges are used, ensure that they (including the pouch intact) are at room temperature (15-30°C, 59-86°F) for at least 30 minutes before using them for analysis on the Sampinute[™] Analyzer. Analyzing the cartridge before the 30-minute point of room temperature incubation period, may yield false results. Therefore, the user should never open the aluminum pouch, exposing the test cartridge to the ambient environment until ready for immediate use.

LIMITATIONS

- The contents of this kit are to be used for the qualitative detection of SARS-CoV-2 spike proteins from a NPS.
- This test detects both viable (live) and non-viable SARS-CoV-2. Test performance depends on the amount of virus (antigens) in the sample and may or may not correlate with viral culture results performed on the same sample.
- A negative test result may occur if the level of antigens in a sample is below the detection limit of the test or if the sample was collected or transported improperly.
- Failure to follow the Test Procedure may adversely affect test performance and/or invalidate the test result.
- Test results must be evaluated in conjunction with other clinical data available to the physician.
- Positive test results do not rule out co-infections with other pathogens.
- Negative test results are not intended to rule in other non-SARS-CoV-2 viral or bacterial infections.
- Negative results should be treated as presumptive and confirmed with an FDA-authorized molecular assay, if necessary, for clinical management including infection control.
- Clinical performance was evaluated with frozen samples, and test performance may be different with fresh samples.
- If the differentiation of specific coronaviruses and strains is needed, additional testing, in consultation with state or local public health departments, is required.

CONDITIONS OF AUTHORIZATION FOR THE LABORATORY

To assist clinical laboratories using the Sampinute[™] COVID-19 Antigen MIA ("your product" in the conditions below), the relevant Conditions of Authorization are listed below:

- Authorized laboratories using your product will include, with test result reports, all Fact Sheets. Under exigent circumstances, other appropriate methods for disseminating these Fact Sheets may be used, which may include mass media.
- Authorized laboratories using your product will use your product as outlined in the "Sampinute[™] COVID-19 Antigen MIA" Instructions for Use. Deviations from the authorized procedures, including the authorized instruments, authorized clinical specimen types, authorized control materials, authorized other ancillary reagents and authorized materials required to use your product are not permitted.
- Authorized laboratories that receive your product will notify the relevant public health authorities of their intent to run your product prior to initiating tests.
- Authorized laboratories using your product will have a process in place for reporting test

results to healthcare providers and relevant public health authorities, as appropriate.

- Authorized laboratories will collect information on the performance of your product and report to DMD/OHT7-OIR/OPEQ/CDRH (via email: CDRH-EUA-Reporting@fda.hhs.gov), BBB Inc. (via website: www.bbbtech.com) and Celltrion USA, Inc. (via email: Sampinute@celltrion.com or via phone: (201) 988-4615) any suspected occurrence of false positive or false negative results and significant deviations from the established performance characteristics of your product of which they become aware.
- All operators using your product must be appropriately trained in performing and interpreting the results of your product. Use appropriate personal protective equipment when handling this kit, and use your product in accordance with the labeling.
- BBB Inc., Celltrion USA, Inc., and authorized laboratories using your product will ensure that any records associated with this EUA are maintained until otherwise notified by the FDA. Such records will be made available to the FDA for inspection upon request.

CLINICAL PERFORMANCE

Positive clinical specimens in viral transport media (VTM) were used for clinical validation.

The clinical performance of Sampinute[™] COVID-19 Antigen MIA was established with a study using NPS specimens, which have been previously characterized and were supplied frozen by a biorepository in the United States.

In the clinical performance, seventy-two (72) samples were measured, resulting in a sensitivity of 94.4% (34/36) and a specificity of 100.0% (36/36).

	Results of Reference Device (RT- PCR) Abbott RealTi <i>me</i> SARS-CoV-2 assay					95%	CI	
		POS	NEG	Total	Sensitivity	94.4%	80.0%	99.0%
Sampinute™ COVID-19 Antigen MIA	POS	34	0	34	Specificity	100.0%	88.0%	100.0%
	NEG	2	36	38	PPV	100.0%	89.9%	100.0%
	Total	36	36	72	NPV	94.7%	82.7%	98.5%
				Prevalence	50.0%			
				%	97.2%			

Hypothetical Positive and Negative Predictive Values

	PPV			NPV		
Prevalence (%)	Estimates (%)	95% CI (%)		Estimates (%)	95% CI (%)	
1.0	100.0	15.0	100.0	99.9	94.8	100.0
2.0	100.0	26.2	100.0	99.9	94.6	100.0
5.0	100.0	47.0	100.0	99.7	94.2	100.0
10.0	100.0	63.9	100.0	99.4	93.3	100.0
15.0	100.0	72.6	100.0	99.0	92.4	99.9
20.0	100.0	78.0	100.0	98.6	91.5	99.8
25.0	100.0	81.6	100.0	98.2	90.4	99.7
30.0	100.0	84.2	100.0	97.7	89.2	99.5
35.0	100.0	86.1	100.0	97.1	87.9	99.4
40.0	100.0	87.6	100.0	96.4	86.4	99.1
45.0	100.0	88.9	100.0	95.7	84.7	98.9
50.0	100.0	89.9	100.0	94.7	82.7	98.5

Sampinute[™] COVID-19 Antigen MIA

ANALYTICAL PERFORMANCE

a) Limit of Detection

Materials: We conducted the testing with SARS-CoV-2 strain isolated from positive NPS specimens obtained from a biorepository (with a determined titer of 1.2×10^4 TCID₅₀/mL). The specimen was used as the starting point for preparing serial dilution samples. The material was supplied frozen.

The purpose of the LoD studies is to determine the lowest detectable concentration of SARS-CoV-2. The LoD was determined by limiting dilution studies. Dilutions were carried out with reagent solutions. For each test, 50 μ L of the diluted sample was added to a sterile nasal swab before conducting the assay based on the Instruction For Use of the SampinuteTM COVID-19 Antigen MIA test cartridge.

The test consisted of two steps for LoD determination:

1. Tentative LoD Confirmation

The cartridges were first tested in a series of 10-fold dilutions (n=3) to determine the dilution concentration that produced a 100% detection rate (3/3) with the subsequent dilution producing a detection rate of less than 100%.

Based on this testing, the concentration chosen was 1.2×10^2 TCID₅₀/mL.

2. LoD Confirmation

A total of forty (40) replicates were then further tested in a series of 2-fold dilutions (twenty

(20) replicates with two reagent lots) to determine the minimum dilution concentration at which at least 95% of the true positive specimens were tested positive for a given target. 1.2×10^2 TCID₅₀/mL chosen from the "Tentative LoD Confirmation" step was used as the starting point for the dilution.

Based on this testing, the LoD was concluded to be $3.0 \times 10^1 \text{ TCID}_{50}/\text{mL}$.

b) Cross-reactivity and c) Microbial Interference

Cross-reactivity of the cartridges was evaluated by testing various viruses (17) and microorganisms (9 including pooled human nasal wash) that potentially may cross-react with the Sampinute[™] COVID-19 Antigen MIA. The final concentration of each organism is documented in the table below. Each microorganism and virus was prepared in the absence and presence of SARS-CoV-2 at 3xLoD concentration. Both the cross-reactivity and microbial interference studies were conducted in triplicate.

Organism	Final Concentration Tested	Cross-Reactivity Results (Count)	Interference Result (Count)
Coronavirus 229E	5.44 × 10⁵ TCID₅₀/mL	0/3	3/3
Coronavirus OC43	2.91 × 10 ⁵ TCID₅₀/mL	0/3	3/3
Coronavirus NL63	7.05 × 10⁵ TCID₅₀/mL	0/3	3/3
SARS-CoV-1	1.00 × 10 ⁵ PFU/mL	0/3	3/3
MERS	1.02 × 10 ⁵ PFU/mL	0/3	3/3
Adenovirus type 1	1.10 × 10 ⁶ TCID ₅₀ /mL	0/3	3/3
Parainfluenza virus 1	1.13 × 10 ⁶ PFU/mL	0/3	3/3
Parainfluenza virus 2	1.18 × 10 ⁶ PFU/mL	0/3	3/3
Parainfluenza virus 3	1.51 × 10 ⁶ PFU/mL	0/3	3/3
Parainfluenza virus 4	5.42 × 10⁵ PFU/mL	0/3	3/3
Enterovirus	5.80 × 10⁵ PFU/mL	0/3	3/3
Rhinovirus	3.19 × 10⁵ PFU/mL	0/3	3/3
Respiratory Syncytial virus A	6.10 × 10⁵ PFU/mL	0/3	3/3
Human metapneumovirus	1.71 × 10⁵ PFU/mL	0/3	3/3
Coronavirus HKU1	4.07 × 10⁵ PFU/mL	0/3	3/3
Influenza A	1.09 × 10 ⁶ TCID ₅₀ /mL	0/3	3/3
Influenza B 1.09 × 10 ⁶ TCID ₅		0/3	3/3

Cross-Reactivity: Sampinute[™] COVID-19 Antigen MIA - Wet Testing

Candida albicans	3.23 × 10 ⁶ CFU/mL	0/3	3/3
Chlamydia pneumoniae	1.61 × 10 ⁶ IFU/mL	0/3	3/3
Haemophilus influenzae	2.07 × 10 ⁶ CFU/mL	0/3	3/3
Pooled human nasal wash	100%	0/3	3/3
Mycoplasma pneumoniae	1.05 × 10 ⁶ CFU/mL	0/3	3/3
Streptococcus pneumoniae	1.05 × 10 ⁶ CFU/mL	0/3	3/3
Bordetella pertussis	2.55 × 10 ⁶ CFU/mL	0/3	3/3
Legionella pneumophila	3 × 10 ⁶ CFU/mL	0/3	3/3
Streptococcus pyogenes	2 × 10 ⁶ CFU/mL	0/3	3/3

* Testing was performed in triplicate.

** Each microorganism was diluted in BD[™] Universal Viral Transport System.

The results show neither observed cross-reactivity nor microbial interference with the organisms at the concentrations tested.

Of all the tests recommended for cross-reactivity, the remaining two that were not included in the wet testing were analyzed *in silico via* Basic Local Alignment Search Tool managed by National Center for Biotechnology Information to determine the likelihood of leading to cross-reactivity.

• *Pneumocystis jirovecii*: 45.4% homology was found for one particular segment of sequence across 9 % of the sequence. Thus, a very low likelihood of cross-reactivity exists between the pathogens.

• *Mycobacterium tuberculosis*: No sequence homology was found between SARS-CoV-2 and *M. tuberculosis*. Thus, no cross-reactivity exists between SARS-CoV-2.

d) Endogenous interference study

A total of 17 potentially interfering substances, either naturally present in respiratory specimens or artificially introduced into the nasal cavity or nasopharynx, were tested in this study to evaluate the susceptibility of the Sampinute[™] COVID-19 Antigen MIA test cartridges to potentially interfering substances when elevated levels of these substances were added to

SARS-CoV-2 positive or negative samples.

The potentially interfering substances were spiked into the SARS-CoV-2 positive or negative samples at elevated levels. For each test, a sterile nasal swab was swirled in the diluted samples before conducting the assay based on the Instruction For Use of the Sampinute[™] COVID-19 Antigen MIA test cartridge.

Concentrations of potentially interfering substances were tested and the study results are summarized in the table below.

Substance	Concentration	SARS-CoV-2 Positive Sample	SARS-CoV-2 Negative Sample	
Whole Blood	4%	3/3	0/3	
Mucin (Bovine submaxillary gland)	0.5%	3/3	0/3	
Tamiflu (Oseltamivir)	5 mg/mL	3/3	0/3	
Mupirocin	10 mg/mL	3/3	0/3	
Tobramycin	4 μg/mL	3/3	0/3	
Sore Throat Phenol Spray	15% v/v	3/3	0/3	
Homeopathic Nasal Spray (Alkalol)	1:10 dilution	3/3	0/3	
Ricola (Menthol)	1.5 mg/mL	3/3	0/3	
Sucrets (Dyclonine/Menthol)	1.5 mg/mL	3/3	0/3	
Chloraseptic (Menthol/Benzocaine)	1.5 mg/mL	3/3	0/3	
CVS Nasal Drops (Phenylephrine) 15% v/v		3/3	0/3	
Afrin (Oxymetazoline) 15% v/v		3/3	0/3	

Potentially Interfering Substances Study: Sampinute[™] COVID-19 Antigen MIA

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CVS Nasal Spray (Cromolyn)	15% v/v	3/3	0/3
Naso GEL (NeilMed)	5% v/v	3/3	0/3
Zicam	5% v/v	3/3	0/3
Fisherman's Friend	1.5 mg/mL	3/3	0/3
Fluticasone Propionate	5% v/v	3/3	0/3

The study results show no interference with the Sampinute[™] Analyzer and the Sampinute[™] COVID-19 Antigen MIA test cartridges observed in the presence of potentially interfering substances at the concentrations tested in this study.

e) High-dose hook effect

No high-dose hook effect was observed up to 3 × 10⁵ TCID₅₀/mL of SARS-CoV-2 when measured with the Sampinute[™] Analyzer and Sampinute[™] COVID-19 Antigen MIA.

f) Matrix equivalency

SARS-CoV-2 strain was formulated in four (4) negative matrices: BD universal viral transport system (UVT), COPAN universal transport medium (UTM), Centers for Disease Control and Prevention VTM (DSR-052-02), and the reagent solution with less than 0.1% sodium azide (contained in Reagent Tube) at a final concentration of 1 x LoD.

The transport media systems containing the contrived samples were stored at 2 - 8°C (36 - 46°F). Each transport media was tested in three (3) replicates and 3 Lots.

All four (4) matrices (BD UVT, COPAN UTM, VTM made according to the CDC SOP (DSR-052-02), and the reagent solution with less than 0.1% sodium azide) show the same results with %CV of less than 10%.

ASSISTANCE

If you have any questions regarding the use of this product or if you want to report a test system problem, please contact BBB Inc. (*via* website: <u>www.bbbtech.com</u>) or Celltrion USA, Inc. (*via* email: Sampinute@celltrion.com, or *via* phone: (201) 988-4615). Test system problems may also be reported to the FDA through the MedWatch medical products reporting program (phone: 800.FDA.1088; fax: 800.FDA.0178; http://www.fda.gov/medwatch).

SYMBOLS



Sampinute[™] COVID-19 Antigen MIA

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