





COV-19C25

# COVID-19 Antigen Rapid Test Device

Kit Contents

- 25 Test Cassettes
- Buffer Nozzle
- Individually packed swabs
- Tube and tube stand
- Product Insert

For the direct and qualitative detection of SARS-CoV-2 viral nucleoprotein antigens from nasopharyngeal secretions and Oropharyngeal secretions





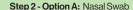






# Procedure Card COVID-19 Antigen Rapid Test Device







Step 2a
Tilt patient's head 70°. Insert the swab
at least 0.5 inch inside the nostril
(nares) until mild resistance was
encountered at the middle turbinate.



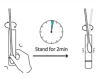
**Step 2a.1**Using a circular motion, the nasal orifice should be swabbed for a minimum of five seconds.



Step 2a.2 Compress the nostril with the fingers to trap the swab tip and rotate the tip for a minimum of five seconds.



Step 2a.3 Repeat steps for other nostril.



Step 3
Insert the swab into the extraction tube. Mix well and squeeze the swab 10-15 times. After waiting 2 minutes, twist the swab against the inner wall as your remove it, trying to release as much liquid as you can while removing.



Step 1 Gently mix the extraction buffer.

Add 10 drops into the extraction tube.

### Step 2 - Option B: Nasopharyngeal Swab



Tilt patient's head 70°. Insert the swab into the nostril parallel to the palate, and gently push the swab into the posterior nasopharynx.

Rotate against nasal well to ensure swab contains cells and mucus.



Insert the nozzle into the sample extraction tube.



Step 5
Add 3 drops of the solution into the sample well by gently squeezing the tube.

### Step 2 - Option C: Oropharyngeal Swab

### Step 2c

Insert the swab completely from the mouth into the throat, centering in on the red part of the throat wall and maxillary tonsils. Rub the bilateral throat tonsils and throat wall with moderate pressure.

Avoid touching the tongue and remove the swab.



Step 6 Read results after 15 minutes.

### **Result Interpretations**



ositive Negative

Invalid

The Rapid Response™ COVID-19 Antigen Rapid Test Device is an in vitro immunochromatographic assay for the direct and qualitative detection of SARS-CoV-2 viral nucleoprotein antigens from nasal and nasopharyngeal secretions from individuals suspected of COVID-19 within 6 days of symptom onset. This test is authorized for use at the Point of Care i.e., in patient care settings.

Results are for the identification of SARS-CoV-2 viral nucleoprotein antigen. Antigens are generally detectable in nasopharyngeal and nasal secretions during the acute phase of infection. Positive results indicate the presence of viral antigens, but clinical correlation with patient history and other diagnostic information is 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 are required to report all positive results to the appropriate public health authorities.

Negative results should be treated as presumptive, and do not rule out SARS-CoV-2 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 the presence of clinical signs and symptoms consistent with COVID-19, and confirmed with a molecular assay, if necessary, for patient

The Rapid Response<sup>TM</sup> COVID-19 Antigen Rapid Test Device is intended for use by trained laboratory personnel or health care professionals.

### PRINCIPLE

The Rapid Response<sup>TM</sup> COVID-19 Antigen Rapid Test Device detects SARS-CoV-2 viral antigens through visual interpretation of colour development. Anti-SARS-CoV-2 antibodies are immobilized on the test region of the nitrocellulose membrane. Anti-SARS-CoV-2 antibodies conjugated to coloured particles are immobilized on the conjugated pad. A sample is added to the extraction buffer which is optimized to release the SARS-CoV-2 antigens

During testing, the extracted antigens bind to anti-SARS-CoV-2 antibodies conjugated to coloured particles. As the specimen migrates along the strip by capillary action and interacts with reagents on the membrane, the complex will be captured by the anti-SARS-CoV-2 antibodies at the test region. Excess coloured particles are captured at the internal control

The presence of a coloured band in the test region indicates a positive result for the SARS-CoV-2 viral antigens, while its absence indicates a negative result. A coloured band at the control region serves as a procedural control, indicating that the proper volume of specimen has been added and membrane wicking is working.

### MATERIALS

Extraction buffer

Nozzle with filter

Tube stand

### Materials Provided

- Individually packed test devices
- Extraction tube
- Individually packed swabs
- Package insert

Materials Required but Not provided

Clock, timer, or stopwatch Materials provided upon request

COVID-19 Antigen Controls: Positive and Negati

### PRECAUTIONS

- For in vitro Diagnostic Use Only
- Read the Product Insert prior to use. Directions should be read and followed carefully. Do not use kit or components beyond the expiration date.
- The device contains material of animal origin and should be handled as a potential biohazard. Do not use if pouch is damaged or open.
- Test devices are packaged in foil pouches that exclude moisture during storage. Inspect each foil pouch before opening. Do not use devices that have holes in the foil or where the pouch has not been completely sealed. Erroneous result may occur if test reagents or components are improperly stored.
- Do not use the Extraction Buffer if it is discoloured or turbid. Discolouration or turbidity may be a sign of microbial contamination.
- All patient specimens should be handled and discarded as if they are biologically hazardous. All specimens must be mixed thoroughly before testing to ensure a representative sample prior to testing.
- Failure to bring specimens and reagents to room temperature before testing may decrease assay sensitivity. Inaccurate or inappropriate specimen collection, storage, and transport may yield false negative test results.
- Avoid skin contact with buffer.
- If infection with SARS-CoV-2 is suspected based on current clinical and epidemiological screening criteria recommended by public health authorities, specimens should be collected with appropriate infection control precautions and sent to state or local health departments for testing
- Viral isolation in cell culture and initial characterization of viral agents recovered in cultures of SARS-CoV-2 specimens are NOT recommended, except in a BSL3 laboratory using BSL3 work practices.

# STORAGE AND STABILITY

- Store the Rapid Response<sup>TM</sup> COVID-19 Antigen Rapid Test at 2~30°C when not in use.
- DO NOT FREEZE
- Kit contents are stable until the expiration dates marked on their outer packaging and

### SPECIMEN COLLECTION AND STORAGE

### Nasal swab (N swab):

- Remove the swab from its packing.
   Tilt patient's head back 70°. Insert the swab through the anterior nares in contact with nasal septum at least 0.5 inches inside the nostril until mild resistance is encountered at the middle turbinate.
- 3) Using a circular motion, the nasal orifice should be swabbed for a minimum of five seconds.
- 4) Compress the nostril with the fingers to trap the swab tip and rotate the tip for a minimum of five seconds.
- 5) Remove and repeat for the other nostril with the same swab.6) Process the swab as soon as possible after collecting the specimen.

# Nasopharyngeal swab (NP swab):

- Remove the swab from its packing.
   Gently insert the sterile swab into the nostril parallel to the palate, not upwards. The distance should be equivalent to that from the ear to the nostril of the patient, indicating contact with the nasopharynx, or until resistance is encountered. Gently rub and roll the swab, leave in place several seconds to saturate tip with secretions. Slowly remove the swab while rotating it.
- 3) Process the swab as soon as possible after collecting the specimen

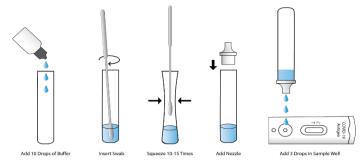
- 1. Use only synthetic fiber swabs with plastic shafts. Do not use calcium alginate swabs or swabs with wooden shafts, as they may contain substances that inactivate some viruses and inhibit further testing.
- Swab specimens should be tested as soon as possible after collection. Use freshly collected specimens for best test performance
- 3. If not tested immediately, swab specimens may be stored at 2-8°C for 21 hours after collection.
- 4. Do not use specimens that are obviously contaminate with blood, as it may interfere with the flow of sample and with the interpretation of test results.

# TEST PROCEDURE

Bring devices, reagents, and specimens and/or controls to room temperature (15~30°C) before

- 1. Label a clean extraction tube with patient or control identification and place it into the tube
- 2. Gently mix extraction buffer. Without touching the buffer bottle to the extraction tube, add 10 drops into the extraction tube.
- 3. Insert the swab with the collected specimen into the extraction tube. Swirl the swab, mixing well. Squeeze the swab 10-15 times by compressing the walls of the tube against the swab.
- 4 Let the solution stand for 2 minutes.
- 5. Remove the swab while pressing the swab head firmly against the inner wall of the tube to release as much liquid as possible. Dispose of the used swab in accordance with the appropriate biohazard waste disposal protocol.
- 6. For each specimen, open the foil pouch just before testing and remove the test device and put it on a clean, level surface. For best results, the assay should be performed within one hour. Label the test device with patient or control identification
- 7. Attach nozzle to sample extraction tube. Invert the tube and add 3 drops of the extracted

solution into the sample well of the test device by gently squeezing the tube. 8. Start the timer. Wait for coloured line(s) to appear. Read results at 15 minu



### RESULT INTERPRETATION



**POSITIVE:** Two coloured bands appear on the membrane. One band appears in the control region (C) and another band appears in the test



No apparent coloured band appears in the test region (T). INVALID: Control band fails to appear. Results from any test which has not produced a control band at the specified read time must be discarded. Please review the procedure and repeat with a new test. If the problem

persists, discontinue using the kit immediately and contact your local

**NEGATIVE:** Only one coloured band appears, in the control region (C).

### NOTE:

- The colour intensity in the test region (T) may vary depending on the concentration of analytes present in the specimen. Therefore, any shade of colour in the test region should be considered positive. Note that this is a qualitative test only and cannot determine the concentration of analytes in the specimen.
- Insufficient specimen volume, incorrect operating procedure or expired tests are the most likely reasons for control band failure.

### QUALITY CONTROL

**Internal Procedural Controls** The Rapid Response<sup>TM</sup> COVID-19 Antigen Rapid Test has built-in (procedural) controls Each test device has an internal standard zone to ensure proper sample flow. The user should confirm that the coloured band located at the "C" region is present before reading the result.

### **External Positive and Negative Controls**

distributor

Good laboratory practice suggests that positive and negative external controls are run routinely to ensure that the test is correctly performed. External positive and negative controls should be used in accordance with applicable accrediting organizations. However, BTNX recommends that labs receiving this test execute a control test for each lot of kits that

### **COVID-19 Antigen Controls:**

Positive and Negative controls are provided upon request with the kit. These controls should be used according to the nasopharyngeal swab test procedure provided in this package insert.

### LIMITATIONS OF THE TEST

- The Rapid Response  $^{TM}$  COVID-19 Antigen Rapid Test Device is for professional invitro diagnostic use and should only be used for the qualitative detection of SARS-CoV-2 antigen. The intensity of colour in a positive band should not be evaluated as "quantitative or semi-quantitative".
- Both viable and nonviable SARS-CoV-2 viruses are detectable with the Rapid Response<sup>TM</sup> COVID-19 Antigen Rapid Test Device.

  As with all diagnostic tests, a definitive clinical diagnosis should not be based on the
- results of a single test but should only be made by the physician after all clinical and laboratory findings have been evaluated.
- laboratory findings have been evaluated. Failure to follow the test procedure and result interpretation may adversely affect test performance and/or invalidate the test result.

  Results obtained with this assay, particularly in the case of weak test lines that are difficult to interpret, should be used in conjunction with other clinical information available to the physician.
- Negative results do not preclude SARS-CoV-2 infection and should be confirmed via molecular assay.
- The performance of the device has not been assessed on specimens from individuals who have been infected with emerging variants of SARS-CoV-2 of public health
- The performance of this device has not been assessed in a population vaccinated against COVID-19.
- This assay is not intended for home testing (or self-testing).

### PERFORMANCE CHARACTERISTICS

Analytical Sensitivity (Limit of Detection): The limit of detection was determined with a quantified SARS-CoV-2 virus and has been evaluated at  $2\times10^{2.4}$  TCID<sub>50</sub>/mL.

## Clinical Evaluation:

Study 1: With Nasal Swab as a sample type: Clinical performance characteristics for Rapid Response<sup>TM</sup> COVID-19 Antigen Rapid Test Device were evaluated in a multi-site prospective, single-blind, controlled clinical trial in the USA from October 2020 to December 2020. The study was performed by Point-of-Care operators with no laboratory experience. These clinical trials were aimed to evaluate the operators with no haboratory experience. These clinical trials were aimed to evaluate the performance of the Rapid Response<sup>™</sup> COVID-19 Antigen Rapid Test Device with a Nasal Swab by comparing with an FDA EUA approved RT-PCR comparator.

The performance of the Rapid Response<sup>™</sup> COVID-19 Antigen Rapid Test Device was established with direct anterior nasal swabs collected from individual symptomatic patients

(within 7 days from symptom onset) who were suspected of COVID-19. Two samples from each patient were collected – one for PCR and another for the Rapid Antigen test. 51 positive specimens and 128 negative specimens were confirmed by RT-PCR.

Table 1: Rapid Response<sup>™</sup> COVID-19 Antigen Rapid Test Clinical Evaluation with Nasal Swabs:

		RT-PCR		Total	
		Positive	Negative	Total	
Rapid Response <sup>TM</sup> COVID-19 Antigen Rapid	Positive	46	0	46	
Test Device	Negative	5	128	133	
Total		51	128	179	
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Diagnostic Sensitivity: 90.2% (78.6% ~ 96.2%)\* Diagnostic Specificity: 100.0% (96.5% ~ 100.0%)\* Overall Agreement: 97.2% (93.2% ~ 98.9%)\* \*95% Confidence Interval

Study 2: Clinical trials for the Rapid Response<sup>TM</sup> COVID-19 Antigen Rapid Test Device were performed at two Point-of-care sites in the USA from August 2020 to October 2020. These clinical trials were aimed to evaluate the performance of the Rapid Response™ COVID-19 Antigen Rapid Test Device by comparing with an RT-PCR comparator.

The performance of the Rapid Response™ COVID-19 Antigen Rapid Test Device was

established with 82 direct nasopharyngeal swabs collected and enrolled from individual symptomatic patients who were suspected of COVID-19. Samples were freshly collected from 2 sites where the operators were minimally trained. 46 positive specimens and 36 negative specimens were confirmed by RT-PCR.

The performance of the Rapid Response<sup>TM</sup> COVID-19 Antigen Rapid Test Device based on the results from these two sites is summarized below

Table 2: Site 1 for Nasopharvngeal Swab Specimen vs. RT-PCR

		RT-PCR		Total
		Positive	Negative	Total
Rapid Response <sup>TM</sup> COVID-19 Antigen	Positive	23	0	23
Rapid Test Device	Negative	1	21	22
		24	21	45

Diagnostic Sensitivity: 95.8% (79.8% ~ 99.3%)\* Diagnostic Specificity: 100.0% (84.5% ~ 100.0%)\* Overall Agreement: 97.8% (88.4% ~ 99.6%)\*
\*95% Confidence Interval

### Table 3: Site 2 for Nasopharyngeal Swab Specimen vs. RT-PCR

		RT-PCR		Total
		Positive	Negative	Total
Rapid Response <sup>TM</sup> COVID-19	Positive	21	0	21
Antigen Rapid Test Device	Negative	1	15	16
		22	15	37
Antigen Rapid Test Device	Negative	1 22	15	16 37

Diagnostic Sensitivity: 95.5% (78.2% ~ 99.2%)\*
Diagnostic Specificity: 100.0% (79.6% ~ 100.0%)\* Overall Agreement: 97.3% (86.2% ~ 99.5%)\* \*95% Confidence Interval

### Table 4: Site 1 & 2 Combined for Nasopharyngeal Swab Specimen vs. RT-PCR

		RT-F	Total	
		Positive	Positive Negative	
Rapid Response <sup>TM</sup> COVID-19	Positive	44	0	44
Antigen Rapid Test Device	Negative	2	36	38
		46	36	82

Diagnostic Sensitivity: 95.6 % (85.5% ~ 98.8%)\* Diagnostic Specificity: 100.0 % (90.4% ~ 100.0%)\* Overall Agreement: 97.6 % (91.5% ~ 99.3%)\* \*95% Confidence Interval

### Cross Reactivity:

Cross reactivity with the following organisms has been studied. Samples positive for the following organisms were found negative when tested with the Rapid Response™ COVID-19 Antigen Rapid Test Device.

HCoV-HKU1	Influenza A (H5N1)	Coxsackie virus A16
HCoV-OC43	Influenza A (H7N9)	Norovirus
HCoV-NL63	Influenza A (H7N7)	Mump virus
HCoV-229E	Influenza B Victoria lineage	Legionella pneumophila
Measles virus	Influenza B Yamagata lineage	Mycoplasma pneumoniae
Streptococcus pneumoniae	Respiratory syncytial virus	Chlamydia pneumoniae
Epstein-Barr virus	Adenovirus	Streptococcus pyogenes
Bordetella Para pertussis	Parainfluenza 1/2/3 virus	Streptococcus agalactiae
Influenza A (H1N1) pdm09	Human metapneumovirus	Group C Streptococcus
Influenza A (H3N2)	Rhinovirus	Staphylococcus aureus

### **Microbial Interference Study:**

Potential microbial interference was evaluated to demonstrate that false negatives will not occur when SARS-CoV-2 is present in a specimen with other microorganisms. Low concentration of SARS-CoV-2 (3 X LOD) was spiked into the higher concentrations of interfering organism and it was found that there is no microbial interference for following

organisms.					
HCoV-HKU1	Influenza A (H5N1)	Coxsackie virus A16			
HCoV-OC43	Influenza A (H7N9)	Haemophilus influenzae			
HCoV-NL63	Influenza A (H7N7)	Candida albicans			
HCoV-229E	Influenza B Victoria lineage	Mycobacterium tuberculosis			
Measles virus	Influenza B Yamagata lineage	Norovirus			
Streptococcus pneumoniae	Respiratory syncytial virus	Mump virus			
Epstein-Barr virus	Adenovirus	Legionella pneumophila			
Bordetella Para pertussis	Parainfluenza 1/2/3 virus	Mycoplasma pneumoniae			
Influenza A (H1N1) pdm09	Human metapneumovirus	Chlamydia pneumoniae			
Influenza A (H3N2)	Rhinovirus	Streptococcus pyogenes			
Group C Streptococcus	Staphylococcus aureus	Streptococcus agalactiae			
Pooled human nasal wash – representative of normal respiratory microbial flora					

### Interfering Substances

The following substances, naturally present in respiratory specimens or that may be artificially introduced into the respiratory tract, were evaluated at the concentrations listed below. None of them were found to affect test performance of the Rapid Response<sup>TM</sup>

Substance	Concentration	Substance	Concentration
3 OTC nasal sprays	10%	Guaiacol glyceryl ether	20 mg/ml
3 OTC mouthwashes	10%	Mucin	1%
3 OTC throat drops	10%	Mupirocin	250 μg/ml
4-acetamidophenol	10 mg/ml	Oxymetazoline	10 mg/ml
Acetylsalicylic acid	20 mg/ml	Phenylephrine	10 mg/ml
Albuterol	20 mg/ml	Phenylpropanolamine	20 mg/ml
Chlorpheniramine	5 mg/ml	Relenza®(zanamivir)	20 mg/ml
Dexamethasone	5 mg/ml	Rimantadine	500 ng/ml
Dextromethorphan	10 mg/ml	Tamiflu ® (oseltamivir)	100 mg/ml
Diphenhydramine	5 mg/ml	Tobramycin	40 mg/ml
Doxylamine succinate	1 mg/ml	Triamcinolone	14 mg/ml
Flunisolide	3 mg/ml		

### **High Dose Hook Effect**

To high dose hook effect was observed when tested with up to a concentration of 1 x 10<sup>6,4</sup> TCID<sub>50</sub>/mL of heat inactivated SARS-CoV-2 virus with the Rapid Response<sup>TM</sup> COVID-19

GLOSSARY OF SYMBOLS					
	Consult instructions for use	$\sum$	Tests per Kit	REF	Catalogue number
2°C (86°F)	Store between 2°C to 30°C	$\square$	Use by date	2	Do Not Reuse
IVD	In vitro diagnostic medical device	LOT	Lot Number	EC REP	Authorized Representative



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CE EC REP MDSS GmbH 30175 Hannover, Germany





# COVID-19 Medical Device Authorization with Conditions for Importation or Sale

# Autorisation d'importation ou de mise en vente d'un instrument médical relatif au COVID-19 avec conditions

**Authorization Reference Number:** 321669 **Numéro de référence de l'autorisation** 

Issue Date: 2021-02-19 Date de délivrance:

Device Class/Classe de l'instrument : 4

Pursuant to section 5 of the Interim Order Respecting the Importation and Sale of Medical Devices for Use in Relation to COVID-19, made by the Minister of Health on March 18, 2020, the medical device listed below is now authorized for sale or importation in Canada.

Each shipment of a COVID-19 medical device that is imported into Canada must be accompanied by a copy of this authorization document. Please ensure to highlight the **Authorization reference number** during the import declaration process to facilitate port entry without any delays.

This authorization is only valid for so long as the Interim Order Respecting the Importation and Sale of Medical Devices for Use in Relation to COVID-19 is in effect.

Conformément à l'article 5 de *l'Arrêté* d'urgence concernant l'importation et la vente d'instruments médicaux relatifs au Covid-19, réalisé par la ministre de la Santé le 18 mars 2020, les instruments indiqués cidessous sont présentement autorisés pour la mise en vente ou l'importation au Canada.

Tout envoi d'un instrument médical relatif au COVID-19 doit être accompagné d'une copie de la présente autorisation. Veuillez vous assurez de souligner le **numéro de référence de l'autorisation** durant le processus de déclaration d'importation pour faciliter l'entrée sans délais aux points de contrôle frontalier.

Cette autorisation est uniquement valide tant que *l'Arrêté d'urgence concernant l'importation et la vente d'instruments médicaux relatifs au Covid*-19 est en vigueur, ou l'autorisation est annulée.

### Device Name(s) Nom de l'instrument

### RAPID RESPONSE COVID-19 ANTIGEN RAPID TEST DEVICE

Name & Address of Authorization Holder/Nom & adresse du titulaire de l'autorisation

BTNX INC.
570 HOOD RD, UNITS #23, 21, 13, 6, 3 AND 5
MARKHAM, ONTARIO
CANADA
L3R 4G7

David Boudreau, ing., Director General, Medical Devices Directorate Directeur général, Direction des instruments médicaux



# Components/Parts/Accessories/Devices for this Licence Les composantes, parties, accessoires et instruments médicaux pour cette homologation

### RAPID RESPONSE COVID-19 ANTIGEN RAPID TEST DEVICE

Device ID/No de l'instrument: 1029056 Device Identifier / Identificateur de l'instrument (Model/Catalog Detail/No de modèle/Catalogue): COV-19C25