PROGRAM Standard Operating Procedure – Laboratory Services		
Title: MIC82200 –	Policy Number:	
BioFire Respiratory Panel 2.1		
Program Name: Laboratory Services		
Applicable Domain: Epidemic/Pandemic		
Additional Domain(s): Lab, DI and Pharmacy Services		
Effective Date: Next Review Date:		
Issuing Authority:	Date Approved:	
Director of Health Services		
Accreditation Canada Applicable Standard: N/A		

GUIDING PRINCIPLE:

The BioFire Respiratory Panel 2.1 (RP2.1) is a real-time, nested multiplexed PCR test designed to simultaneously identify nucleic acids from 22 different viruses and bacteria associated with respiratory tract infection, including SARS-CoV-2.

PURPOSE/RATIONALE:

This standard operating procedure describes the BioFire Respiratory Panel 2.1 test using the BioFire Torch at Stanton Territorial Hospital.

SCOPE/APPLICABILITY:

This procedure applies to Medical Laboratory Technologists (MLTs) processing specimens for respiratory pathogens using the BioFire Torch.

SAMPLE INFORMATION:

Туре	Swab
Type	Saline gargle
	Nasopharyngeal
Source	Throat/oropharyngeal
	Saline gargle rinse
Collection	Viral transport media
Collection	Universal transport media
Container	Saline gargle tube
	VTM/UTM:
	Room temperature up to 4 hours
Ctobility	Refrigerated up to 7 days
Stability	• Frozen (\leq -15 °C or \leq -70°C) up to 30 days
	Saline gargle:
	Refrigerated up to 5 days

Storage Requirements	Room temperature or refrigerated
Criteria for	 Unlabeled/mislabeled samples Sample container label does not match patient
rejection	identification on requisition Sample not in UTM/VTM or saline gargle tube Sample not stored correctly

REAGENTS and/or MEDIA:

- BioFire RP2.1 pouch
- Hydration Injection Vial (blue)
- Sample Injection Vial (red)
- Sample Buffer Ampoule
- Pouch loading station
- Accel TB 1 L bottle
- Accel TB wipes
- 70% isopropyl alcohol
- 10% bleach solution

SUPPLIES:

- Personal protective equipment
- Absorbent bench liner
- Sample racks
- Wet waste container
- Dry waste container

- Orange autoclave bag
- Spray bottles
- Transfer pipettes provided in kit
- Sharps container
- Blue pouch tray

EQUIPMENT:

- BioFire Torch
- Printer
- Class II biosafety cabinet
- Vortex mixer
- Refrigerator

ENVIRONMENTAL CONTROLS:

- Store the test kit, including reagent pouches and buffers, at room temperature (15°C to 25°C) DO NOT REFRIGERATE
- Avoid storage of any materials near heating or cooling vents or in direct sunlight
- All kit components should be stored and used together. Do not use components from one kit with those of another kit
- Once the pouch packaging has been opened, the pouch should be loaded with sample as soon as possible (within 30 minutes)
- Once a pouch has been loaded with sample, the test run should be started as soon as possible (within approximately 60 minutes)
- To prevent amplicon contamination: discard used pouches after the run has completed, avoid excessive handling of pouches after test runs and avoid exposing pouches to sharp edges or anything that might cause a puncture

SPECIAL SAFETY PRECAUTIONS:

- Patient samples should only be opened and prepared for testing in a contained environment (i.e., certified Class II BSC)
- Personal Protective Equipment (PPE) required when working with suspect SARS-CoV-2 samples includes:
 - > Lab gown, double layer nitrile gloves and Class II BSC
- The test operator and all personnel in the immediate vicinity should be wearing appropriate PPE at all times when working with suspect respiratory pathogen samples, in the event of a spill outside of the BSC
- All items in the BSC must be sprayed or wiped with 70% isopropyl alcohol or Accel TB and allowed to disinfect for 5 minutes prior to removal
- All personnel handing potential respiratory pathogen samples should be knowledgeable in their laboratory's biological spill clean-up protocol for infectious respiratory samples
- Handle all samples and waste materials as if they were capable of transmitting infectious agents

QUALITY CONTROL:

- Refer to MIC82300-BioFire RP2.1 Quality Control for quality control procedure
- Record all results on MIC82310-BioFire RP2.1 Quality Control Results Record

PROCEDURE INSTRUCTIONS:

Step	Action
Prepa	ring the BSC
1	 Order BioFire RP2.1 testing in the LIS: In SoftMic, accession the order using the test code PCRES NOTE: Make sure the priority is STAT Place the sample barcode label and media barcode label in the pouch of the biohazard bag Place samples in the blue bin labelled PCRES in the microbiology specimen fridge
2	Ensure the daily maintenance for the BioFire has been completed and documented on MIC82110-Maintenance Record-BioFire.
3	 Set up the clean BSC with the following: Absorbent pad on working surface Sample racks Wet waste container half full with Accel TB Dry waste container containing an autoclave bag Sharps container Spray bottles with 70% isopropyl alcohol and with Accel TB Accel TB wipes with top opened Clean blue tray Pouch Loading Station BioFire Respiratory Panel 2.1 pouch, Sample Buffer Ampoule, Hydration Injection Vial, Sample Injection Vial, and transfer pipette

Step	Action	
Prepa	Preparing the Pouch	
1	In the BSC, open biohazard bag and discard in the dry waste container. Wipe or spray each sample with Accel TB and place in the sample rack. Once dry, label the sample with the sample label. Leave the media barcode label on the right-hand side of the working area.	
2	Apply the media barcode label to the tab of the pouch NOTE: Do not cover the barcode label on the front of the pouch	
3	Insert the pouch into the Pouch Loading Station, aligning the red and blue labels on the pouch with the red and blue arrows on the Pouch Loading Station.	
4	Place a red-capped Sample Injection Vial into the red well of the loading station. Place a blue-capped Hydration Injection Vial into the blue well of the Pouch Loading Station.	

Step	Action
Hydra	ting the Pouch
1	Unscrew the Hydration Injection Vial from the blue cap.
2	Insert the Hydration Injection Vial's cannula tip into the Hydration Port located directly below the blue arrow of the Pouch Loading Station.
3	Forcefully push down in a firm and quick motion to puncture the seal until a faint "pop" is heard and there is an ease in resistance. Wait as the correct volume of hydration solution is pulled into the pouch by vacuum.
4	 Verify that the pouch has been hydrated: Flip the barcode label down and check to see that fluid has entered the reagent wells (located at the base of the rigid plastic part of the pouch). Small bubbles may be seen. Loss of volume in injection vial Hydration fluid in last 11 wells

Step	Action
Preparing the Sample Mix	
1	 Add Sample Buffer to the Sample Injection Vial: Hold the Sample Buffer Ampoule with the tip facing up Firmly pinch at textured plastic tab on the side of the ampoule until the seal snaps

2	Invert the ampoule over the red-capped Sample Injection Vial and dispense the Sample Buffer using a slow forceful squeeze. Repeat this with a second squeeze. NOTE: Avoid generating foam
3	Thoroughly mix the sample with the vortex mixer for 10 seconds.
4	Open the sample carefully to avoid touching droplets on the inner lid and place upright on the absorbent pad to avoid droplets falling on the work surface.
5	Use the transfer pipette provided to draw specimen to the third line (approximately 0.3 mL) on the transfer pipette.
6	Add the specimen to the Sample Injection Vial.
7	Rinse the pipette in the wet waste container with Accel TB and allow to soak for at least 30 minutes.
8	Recap the patient sample tube and move to the back of the rack.
9	Tightly close the lid of the Sample Injection Vial and invert the vial 5 times to mix.
10	Return the Sample Injection Vial to the red well of the loading station.

Step	Action	
Loadi	Loading the Sample Mix	
1	Unscrew the Sample Injection Vial from the red cap and wait for 5 seconds with the vial resting in the cap. NOTE: Waiting 5 seconds decreases the risk of dripping and contamination from the sample	
2	Insert the Sample Injection Vial's cannula tip into the Sample Port located directly below the red arrow of the Pouch Loading Station.	
3	Forcefully push down in a firm and quick motion to puncture seal until a faint "pop" is heard and sample is pulled into the pouch by vacuum.	
4	 Verify that the sample has been loaded: Flip the barcode label down and check to see that fluid has entered the reagent well next to the sample loading port If the pouch fails to pull sample from the Sample Injection Vial, the pouch should be discarded and begin again with a new pouch 	

	Loss of volume in injection vial
	Sample drawn into 1 st well
5	Discard the Sample Injection Vial and the Hydration Injection Vial into the sharps container.
6	Remove the pouch from the Pouch Loading Station and place on the left- hand side on the blue tray. NOTE: If any of the sample mix drips onto the Pouch Loading Station remove from use and soak in 10% bleach for 15 minutes before being put into use again
7	Wipe Pouch Loading Station with Accel TB before beginning next sample.
8	Spray outer gloves thoroughly with isopropyl alcohol, rub together and allow to air dry. NOTE: Always disinfect gloves between loading of each sample
9	Repeat pouch loading procedure for up to 15 additional samples.

Step	Action	
Creati	Creating a Test Run	
1	Remove outer gloves and transfer the loaded pouches in the blue tray to the BioFire bench.	
2	On the BioFire Torch dashboard, select an available module.	
3	Using the built-in scanner, scan the pouch barcode and the sample ID barcode. Insert the pouch into the module.	
4	Enter the operator username: micro and the password: micro . Select Next . Review the entered information and select START RUN .	

Step	Action
Cleaning the BSC	
1	Remove gloves and don a new pair.
2	In the BSC, spray samples with Accel TB and place the sample rack on the cart.
3	Wipe the blue tray and "clean" area of the BSC with an Accel TB wipe.
4	Remove gloves and don a new pair. Transfer samples to the fridge.

Step	Action		
Generating a Test Report			
1	A report is generated automatically upon completion of a run.		
2	To view runs or reprint: Select Browse Runs on the main screen. Select the sample \rightarrow View report \rightarrow Print.		

INTERPRETATION OF RESULTS:

RESULT	INTERPRETATION		
Not Detected	The pouch controls were successful (Passed)		
Not Delected	 The assay for the organism was negative 		
Detected	 The pouch controls were successful (Passed) 		
Delected	 The assay for the organism was positive 		
	 The pouch controls were successful (Passed) 		
Equivocal	• The combination of positive and negative assay results for		
	Influenza A were inconclusive		
	The pouch controls or run were not successful		
Invalid	Run status is displayed as: Aborted, Incomplete,		
	Instrument Error or Software Error		

REPORTING INSTRUCTIONS:

BF Results:	
SARS-CoV-2	
Influenza A	
Influenza B	
RSV	Report: NEGATIVE for each target
Rhinovirus/	
Enterovirus	
NOT	
DETECTED	

BF Results: SARS-CoV-2	 Repeat testing on the GeneXpert to determine Ct value Add comment in TCOMM that testing was repeated
DETECTED	Add comment in recommend that testing was repeated

	 <u>GeneXpert results are positive</u>:
	Report: POSITIVE
	• If the Ct value is <30.0 add test ?REFE and finalize with a
	"." for send out to APL for COVID sequencing
	Phone to OCPHO (HPU1) at (867)920 8646
BF Results:	\circ Inform OCPHO if Ct is >38
SARS-CoV-2	Phone result to ordering location
DETECTED	Report will automatically print to OCPHO (HPU1)
\mathbf{C}	Report will automatically print to Stanton IPAC (SIPAC)
GX repeat	if ER or inpatient
testing	Check the home address of the patient. If from
POSITIVE	Nunavut:
	 Phone results to the applicable Nunavut CPHO
	 Copy results to the applicable Nunavut CPHO
	Place sample in the red rack in the micro specimen fridge
	 Place GX printout in folder labelled "Positive GX Printouts"

BF Results: SARS-CoV-2 DETECTED GX repeat testing NEGATIVE	 <u>GeneXpert results are negative</u>: Repeat testing on both GeneXpert and BioFire If repeat results are the same: Report: INDETERMINANT From the keypad add key R to add repeat sample collection comment Phone OCPHO (HPU1) at (867) 920 8646 Inform OCPHO that discrepant results between the GeneXpert and BioFire were obtained Phone result to ordering location Report will automatically print to OCPHO (HPU1) Report will automatically print to Stanton IPAC (SIPAC) if ER or inpatient Check the home address of the patient. If from Nunavut:
BF Results: Influenza A-H1 Influenza A- H1-2009 Influenza A-H3 DETECTED	 Report: POSITIVE for Influenza A Report: POSITIVE for the Flu A subtype detected Add test ?REFE and finalize with a "." for send out to APL for Flu A sequencing Report will automatically print to OCPHO (HPU1) Report will automatically print to Stanton IPAC (SIPAC) if ER or inpatient Check the home address of the patient. If from Nunavut copy results to the applicable Nunavut CPHO Place BF printout in folder labelled "Positive BF Printouts"
BF Results: Influenza A-H1 Influenza A- H1-2009 Influenza A-H3 EQUIVOCAL	 If any of the Influenza A targets are equivocal: Repeat testing on the BioFire If the repeat testing is equivocal: Report: POSITIVE for Influenza A Report: POSITIVE for the Flu A subtype equivocal Add test ?REFE and finalize with a "." for send out to APL for Flu A sequencing Report will automatically print to OCPHO (HPU1) Report will automatically print to Stanton IPAC (SIPAC) if ER or inpatient Check the home address of the patient. If from Nunavut copy results to the applicable Nunavut CPHO Place sample in the red rack in the micro specimen fridge Place BF printout in folder labelled "Positive BF Printouts"

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BF Results: Influenza A DETECTED No subtype detected	 Report: POSITIVE for Influenza A Refer sample to APL for subtype confirmation: From the keypad add key V to add referred out comment Order test code VIRC for send out
BF Results: Influenza B DETECTED	 Report: POSITIVE Report will automatically print to OCPHO (HPU1) Report will automatically print to Stanton IPAC (SIPAC) if ER or inpatient Check the home address of the patient. If from Nunavut copy results to the applicable Nunavut CPHO
BF Results: RSV DETECTED	 Report: POSITIVE Phone result to the ordering location Report will automatically print to OCPHO (HPU1) Report will automatically print to Stanton IPAC (SIPAC) if ER or inpatient Check the home address of the patient. If from Nunavut copy results to the applicable Nunavut CPHO
BF Results: Rhino/Entero DETECTED	 Report: POSITIVE Report will automatically print to OCPHO (HPU1) Place sample in the red rack in the micro specimen fridge Place BF printout in folder labelled "Positive BF Printouts"
BF Results: SARS-CoV-2 Influenza A Influenza B RSV Rhinovirus/ Enterovirus INVALID	 Repeat testing on the BioFire Add comment in TCOMM that testing was repeated If the repeat results are invalid: Report: INVALID From the keypad add key R to add repeat sample collection comment Phone the ordering location and request a new sample be collected

NOTE: All calls are to be documented in the Call Box in SoftMic

NOTE: If ordering location is not available, OCPHO will follow up with the patient. Document attempt in the Call Box

Step	Action		
Completing the Run			
1	Check the Resulting Worklist-Torch to ensure all ordered samples are complete.		
2	In the BSC, with gloved hands, remove the used pipettes from the wet waste container and place into the dry waste container.		

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	Remove the autoclave bag from the dry waste container. Tie and wipe		
3	outside with an Accel TB wipe. Place in the biohazard garbage.		
4	Ensure supplies in BSC are stocked up and sharps container is not full.		
5	Wipe the BSC with an Accel TB wipe. Turn off the blower and lower the		
	sash. Remove gloves and don a fresh pair.		
6	Ensure all used pouches from the BioFire are discarded in the biohazard		
	waste.		
	When all testing of patient samples and disinfection of surfaces is complete,		
7	remove PPE and place in the biohazard waste. Retrieve samples from the		
	refrigerator and place in correct storage location.		

LIMITATIONS:

- 1. The BioFire RP2.1 is a qualitative test and does not provide a quantitative value for the organism(s) in the specimen
- 2. Results from this test must be correlated with clinical history
- 3. The performance of BioFire RP2.1 has not been validated for specimens collected from individuals without signs or symptoms of respiratory infection
- 4. The detection of viral and bacterial nucleic acid is dependent upon proper specimen collection, handling, transportation, storage, and preparation.
- 5. A negative BioFire RP2.1 result does not exclude the possibility of viral or bacterial infection
- 6. If four or more organisms are detected in a specimen, retesting is recommended to confirm the polymicrobial result
- Recent administration of nasal influenza vaccines prior to NPS specimen collection could lead to accurate virus detection by the BioFire RP2.1 of the viruses contained in the vaccine, but would not represent infection by those agents

CROSS-REFERENCES:

- MIC82110-Maintenance Record-BioFire
- MIC82300-BioFire RP2.1 Quality Control
- MIC82310-BioFire RP2.1 Quality Control Results Record

REFERENCES:

- 1. *BioFire Respiratory Panel 2.1 (RP2.1)* Instructions for Use (EUA). BFR0000-8303-01, May 2020
- 2. BioFire FilmArray Torch Operator's Manual, HTFA-PRT-0001-05, March 2020
- 3. Canadian Laboratory Response Network Training: COVID-19 Point of Care Diagnostics. *BioFire FilmArray Torch and the FilmArray Respiratory Panel 2.1* (*RP2.1*).
- 4. Biosafety Advisory Committee. *STHA Biosafety Program Manual*. January 2016

Title: MIC82200-BioFire Respiratory Panel 2.1 Issuing Authority: Director of Health Services Next Review Date:

APPROVAL:

Date

REVISION HISTORY:

REVISION	DATE	Description of Change	REQUESTED BY
1.0	10 Aug 20	Initial Release	L. Steven
2.0	02 Oct 20	Updated to remove "presumptive" from reporting	L. Steven
3.0	23 Dec 20	Updated to reflect positive results report to OCPHO, inform OCPHO when CT \geq 38 and record repeat testing in TCOMM	L. Steven
4.0	05 Feb 21	Updated to include the reporting of Influenza A, Influenza B, RSV and Enterovirus/Rhinovirus	L. Steven
5.0	28 Apr 21	Updated to reflect new Xpert Xpress SARS-CoV-2/Flu/RSV cartridge	L. Steven
6.0	17 Oct 22	Updated to reflect notification of Nunavut patients to Nunavut CPHO and sending positive Flu A samples to APL	L. Steven