<b>PROGRAM Standard Operating Procedure – Laboratory Services</b>			
Title: MIC10300 – Xpert <i>C.difficile</i>	Policy Number:		
Program Name: Laboratory Services			
Applicable Domain: Lab, DI and Pharmacy Services			
Additional Domain(s):			
Effective Date:	Next Review Date:		
Issuing Authority:	Date Approved:		
Director of Health Services			
Accreditation Canada Applicable Standard: N/A			

### **GUIDING PRINCIPLE:**

The Xpert *C.difficile* assay is a qualitative *in vitro* diagnostic test for rapid detection of toxin B gene sequences from unformed (liquid or soft) stool specimens collected from patients suspected of having *Clostridium difficile* infection (CDI). The test utilizes automated real time polymerase chain reaction (PCR) to detect toxin gene sequences associated with toxin producing *C.difficile*.

### **PURPOSE/RATIONALE:**

This standard operating procedure describes the Xpert *C.difficile* test using the GeneXpert Dx System.

### SCOPE/APPLICABILITY:

This procedure applies to Medical Laboratory Technologists (MLTs) and Medical Laboratory Assistants (MLAs) processing specimens for *C.difficile* using the GeneXpert Dx System.

Туре	Stool (Unformed)		
Collection	A Orango ton storilo containor		
Container	Orange top, sterile container		
Stability	Room temperature up to 24 hours		
Stability	Refrigerated up to 5 days		
Storage	Deem temperature or refrigerated		
Requirements	Room temperature or refrigerated		
	1. Unlabeled/mislabeled samples		
Criteria for	2. Sample container label does not match patient		
rejection	identification on requisition		
	3. Sample not in sterile container		

### SAMPLE INFORMATION:

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4.	Sample not stored correctly
5.	Repeat testing on positive samples will not be performed
	within 7 days (Cancellation code:  XCDP)
6.	Repeat testing on negative samples will not be
	performed within 7 days (Cancellation code: IXCDN)
7.	Testing will not be performed on patients <12months old
	(Cancellation code:  XCD1)
8.	Testing for <i>C. difficile</i> Toxin is not performed on formed
	stools (Cancellation code: IXCDT)

#### **REAGENTS** and/or MEDIA:

• Xpert *C.difficile* cartridge

### SUPPLIES:

- Personal protective equipment
- Dry waste container

## **EQUIPMENT:**

- GeneXpert Dx System
- Printer
- Class II biosafety cabinet (BSC)

## **ENVIRONMENTAL CONTROLS:**

- Store Xpert C. difficile assay cartridges upright between 2°C to 28°C
- Do not open a cartridge lid until you are ready to perform testing
- Do not touch the Reaction Tube, always handle the cartridge by its Body

# SPECIAL SAFETY PRECAUTIONS:

Containment Level 2 facilities, equipment, and operational practices for work involving infectious or potential infectious materials or cultures.

- Ensure that appropriate hand hygiene practices be used.
- Lab gown must be worn when performing activities with potential pathogens.
- Gloves must be worn when direct skin contact with infected materials is unavoidable.
- Eye protection must be used when there is a known or potential risk of exposure of splashes.
- All procedures that may produce aerosols, or involve high concentrations or large volumes should be conducted in a biological safety cabinet (BSC).
- The use of needles, syringes and other sharp objects should be strictly limited.

All patient specimens are assumed to be potentially infectious. Routine Practices must be followed. Since viable micro-organisms are used, all cultures must be handled with appropriate precautions. All equipment in contact with cultures should be decontaminated by appropriate methods.

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• Vortex mixer

Sample reagent

Sterile, dry swabs

Disposable transfer pipettes

• Refrigerator

# **QUALITY CONTROL:**

- Refer to MIC60080-Xpert *C.difficile* Quality Control for quality control procedure
- Record all results on MIC60081-QC Results Record-Xpert C.difficile

## **PROCEDURE INSTRUCTIONS:**

Step	Action		
Preparing the Run			
1	<ul> <li>Order GeneXpert <i>C.difficile</i> testing in the LIS:</li> <li>Medipatient the order if required</li> <li>In SoftMic, accession the order using the test code PCCDI</li> <li>Add any "copies to" if required</li> <li>Collect, receive and plate the order</li> <li>Label the requisition with the requisition label and scan into SoftMedia</li> <li>Place the sample barcode label on the sample and the barcode label in the pouch of the biohazard bag for when the sample is being processed</li> </ul>		
2	Ensure the daily maintenance for the GeneXpert has been completed and is documented on MIC81110-Maintenance Record-GeneXpert.		

Step	Action			
Prepa	Preparing the Cartridge			
1	Remove the cartridge and reagent from the package. Acquire a swab and a pipette for each sample being tested.			
2	Apply the media barcode label to the right hand side of the cartridge, near the base. NOTE: Do not cover the barcode label on the front of the cartridge			
3	The test must be started within 30 minutes of adding reagents to the cartridge.			
4	Vortex the sample for 10 seconds to ensure it is evenly mixed.			
5	Briefly place a swab in the unformed stool sample. The swab does not need to be completely saturated: Too With sample Correct amount of sample Too much sample Too much sample			
6	Insert the swab into the vial containing the Sample Reagent.			
7	Hold the swab by the stem near the rim of the vial and push the stem against the edge of the vial to break it. <b>NOTE:</b> Make sure the swab is short enough to allow the cap to close			
8	Close the lid and vortex at high speed for 10 seconds.			
9	Pry open the cartridge lid and open wrapper of the transfer pipette.			

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	Transfer the entire contents of the Sample Reagent to the Sample Chamber of the cartridge:
10	Loading Diamber
11	Firmly snap close the lid to seal the cartridge and place in the cartridge
	Tray.

Step	Action		
Creat	Creating a Test Run		
1	Transfer the loaded cartridges in the cartridge tray to the GeneXpert bench.		
2	Log into the GeneXpert software using the username <b>admin1</b> and the password <b>covid19</b> .		
3	Confirm that all modules are detected by the software and ready for testing.		
4	On the GeneXpert software, click <b>Create Test</b> at the top left.		
5	Using the scanner, scan the sample ID barcode and the cartridge barcode. Select <b>Start Test</b> .		
6	Locate the module with the blinking green light, open the module door and load the cartridge.		
7	Close the module door firmly, it will latch closed.		

Step	Action		
Gener	Generating a Test Report		
1	A report is generated automatically upon completion of a run.		
2	To view runs or reprint: Select <b>View Results</b> on the menu bar. Click <b>Report</b> $\rightarrow$ Check <b>Patient ID</b> $\rightarrow$ Click <b>Preview PDF</b> $\rightarrow$ Click <b>Print</b>		

# **INTERPRETATION OF RESULTS:**

RESULT	INTERPRETATION		
Toxigenic <i>C.difficile</i> NEGATIVE	<ul> <li><i>C.difficile</i> target DNA sequences are not detected</li> <li>Toxins producing <i>C.difficile</i> targets not detected</li> </ul>		
Toxigenic <i>C.difficile</i> POSITIVE	<ul> <li><i>C.difficile</i> target DNA sequences are detected</li> <li>The toxin producing <i>C.difficile</i> target(s) have Cts within the valid range</li> </ul>		
NO RESULT	Presence or absence of <i>C.difficile</i> target DNA cannot be determined		
ERROR/ INVALID	<ul> <li>Presence or absence of <i>C.difficile</i> cannot be determined</li> <li>Toxin producing <i>C.difficile</i> targets-NO RESULT</li> <li>Probe Check: FAIL; one or more probe checks have failed</li> </ul>		

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REPORTING INSTRUCTIONS:			
Toxigenic <i>C.difficile</i> <b>NEGATIVE</b>	<ul> <li>Report: NEGATIVE</li> <li>The following comment will be automatically added to the report: "The Xpert C.difficile assay is a qualitative diagnostic test for detection of C.difficile toxin B gene sequences in a stool sample"</li> </ul>		
Toxigenic <i>C.difficile</i> <b>POSITIVE</b>	<ul> <li>Report: POSITIVE</li> <li>Phone results to patient location         <ul> <li>Document call in the Call Box in SoftMic</li> <li>Report will automatically print to OCPHO (HPU1)</li> </ul> </li> <li>Report will automatically print to Stanton IPAC (SIPAC) and Inuvik IPAC (IIPAC) if ER or inpatient</li> <li>The following comment will be automatically added to the report: "The Xpert C.difficile assay is a qualitative diagnostic test for detection of C.difficile toxin B gene sequences in a stool sample"</li> </ul>		
Toxigenic <i>C.difficile</i> <b>NO RESULT</b>	<ul> <li>Retest the sample with a new cartridge         <ul> <li>Add comment in TCOMM that testing was repeated</li> </ul> </li> <li>If repeat testing is the same:         <ul> <li>Phone the ordering location and request a new sample be collected</li> <li>Report: NO RESULT - Presence of C.diff toxin cannot be determined</li> <li>From the keypad add key R to add repeat sample collection comment</li> </ul> </li> </ul>		

# LIMITATIONS:

- 1. This test detects but does not differentiate the NAP1 (Ribotype 027) strain from other toxigenic strains of *C.difficile*.
- 2. This test targets the *tcdB* gene for Toxin B production. This test will not detect strains of *C.difficile* that do not contain the *tcdB* gene.
- 3. Positive results observed with immunocompromised pediatric patients may reflect asymptomatic carriage of *C.difficile*.
- 4. Results from the Xpert *C.difficile* Assay should be interpreted in conjunction with other laboratory and clinical data available to the clinician.
- 5. Erroneous test results might occur from improper specimen collection, failure to follow the recommended sample collection, handling and storage procedures, technical error, sample mix-up, or because the number of organisms in the specimen is too low to be detected by the test.
- 6. Inhibition of the Xpert *C.difficile* Assay has been observed in the presence of the following substances: Zinc oxide paste and Vagisil cream.
- 7. False negative results may occur when the infecting organism has genomic mutations, insertions, deletions or rearrangements or when performed early in the course of illness.

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### **CROSS-REFERENCES:**

- MIC60080-Xpert *C.difficile* Quality Control
- MIC60081-QC Results Record-Xpert *C.difficile* QC
- MIC81110-Maintenance Record-GeneXpert

#### **REFERENCES:**

- 1. Cepheid GeneXpert. Xpert *C. difficile* Assay Instructions for Use. 300-9680 Rev. F, March 2016
- 2. Cepheid GeneXpert. Dx System User Manual. 301-0045, Rev.C, June 2012

### **APPROVAL:**

Date

#### **REVISION HISTORY:**

REVISION	DATE	Description of Change	REQUESTED BY
1.0	30 Jan 22	Initial Release	L. Steven

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