

<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: Director of Health Services	Date Approved:
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.

**SAMPLE INFORMATION:**

<b>Type</b>	Stool (Unformed)
<b>Collection Container</b>	<ul style="list-style-type: none"> <li>Orange top, sterile container</li> </ul>
<b>Stability</b>	<ul style="list-style-type: none"> <li>Room temperature up to 24 hours</li> <li>Refrigerated up to 5 days</li> </ul>
<b>Storage Requirements</b>	Room temperature or refrigerated
<b>Criteria for rejection</b>	<ol style="list-style-type: none"> <li>Unlabeled/mislabeled samples</li> <li>Sample container label does not match patient identification on requisition</li> <li>Sample not in sterile container</li> </ol>

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	<ol style="list-style-type: none"><li>4. Sample not stored correctly</li><li>5. Repeat testing on positive samples will not be performed within 7 days (Cancellation code:  XCDP)</li><li>6. Repeat testing on negative samples will not be performed within 7 days (Cancellation code: IXCDN)</li><li>7. Testing will not be performed on patients &lt;12months old (Cancellation code:  XCD1)</li><li>8. Testing for <i>C. difficile</i> Toxin is not performed on formed stools (Cancellation code: IXCDT)</li></ol>
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**REAGENTS and/or MEDIA:**

- Xpert *C.difficile* cartridge
- Sample reagent

**SUPPLIES:**

- Personal protective equipment
- Dry waste container
- Sterile, dry swabs
- Disposable transfer pipettes

**EQUIPMENT:**

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

**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.

**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
<b>Preparing the Run</b>	
<b>1</b>	Order GeneXpert <i>C.difficile</i> testing in the LIS: <ul style="list-style-type: none"> <li>• Medipatient the order if required</li> <li>• In SoftMic, accession the order using the test code <b>PCCDI</b></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>
<b>2</b>	Ensure the daily maintenance for the GeneXpert has been completed and is documented on MIC81110-Maintenance Record-GeneXpert.

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

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<b>10</b>	Transfer the entire contents of the Sample Reagent to the Sample Chamber of the cartridge: 
<b>11</b>	Firmly snap close the lid to seal the cartridge and place in the cartridge tray.

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

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

**INTERPRETATION OF RESULTS:**

RESULT	INTERPRETATION
<b>Toxigenic <i>C.difficile</i> NEGATIVE</b>	<ul style="list-style-type: none"> <li>• <i>C.difficile</i> target DNA sequences are not detected</li> <li>• Toxins producing <i>C.difficile</i> targets not detected</li> </ul>
<b>Toxigenic <i>C.difficile</i> POSITIVE</b>	<ul style="list-style-type: none"> <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>
<b>NO RESULT</b>	<ul style="list-style-type: none"> <li>• Presence or absence of <i>C.difficile</i> target DNA cannot be determined</li> </ul>
<b>ERROR/ INVALID</b>	<ul style="list-style-type: none"> <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 style="list-style-type: none"><li>• Report: <b>NEGATIVE</b></li><li>• The following comment will be automatically added to the report: <b>"The Xpert <i>C.difficile</i> assay is a qualitative diagnostic test for detection of <i>C.difficile</i> toxin B gene sequences in a stool sample"</b></li></ul>
Toxigenic <i>C.difficile</i> <b>POSITIVE</b>	<ul style="list-style-type: none"><li>• Report: <b>POSITIVE</b></li><li>• Phone results to patient location<ul style="list-style-type: none"><li>➢ Document call in the Call Box in SoftMic</li></ul></li><li>• Report will automatically print to OCPHO (HPU1)</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: <b>"The Xpert <i>C.difficile</i> assay is a qualitative diagnostic test for detection of <i>C.difficile</i> toxin B gene sequences in a stool sample"</b></li></ul>
Toxigenic <i>C.difficile</i> <b>NO RESULT</b>	<ul style="list-style-type: none"><li>• Retest the sample with a new cartridge<ul style="list-style-type: none"><li>➢ Add comment in TCOMM that testing was repeated</li></ul></li><li>• If repeat testing is the same:<ul style="list-style-type: none"><li>➢ Phone the ordering location and request a new sample be collected</li><li>➢ Report: <b>NO RESULT - Presence of <i>C.diff</i> toxin cannot be determined</b></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:**

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Date

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**REVISION HISTORY:**

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

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