# TRAINING UPDATE

Lab Location:	SGMC & WOMC	Date Distributed:	3/10/22
Department:	Microbiology and Processing	Due Date:	4/10/2022

# **DESCRIPTION OF PROCEDURE REVISION**

Name of procedure:

# *Clostridium difficile* Toxin B PCR using Cepheid GeneXpert® (AHC.M1003 v3)

**Description of change(s):** 

**Section 10.6-** Removed old codes for C diff toxin and replaced with codes for reflex.

Section 10.7- Added steps for reflex testing

This revised SOP will be implemented on April 12, 2022

Document your compliance with this training update by taking the quiz in the MTS system.

Technical SOP			
Title	Clostridium difficile Toxin B PCR usin	g Cephe	id GeneXpert®
Prepared by	Ron Master	Date:	2/18/2019
Owner	Ron Master	Date:	2/18/2019

Laboratory Approval	Local Effective Date:	
Print Name and Title	Signature	Date
Refer to the electronic signature		
page for approval and approval		
dates.		

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#### 1. TEST INFORMATION

Assay	Method/Instrument	Test Code
Cepheid GeneXpert	Real-time Polymerase Chain Reaction	CDPCR
Clostridium difficile PCR	(PCR) Assay / GeneXpert System	CDFCK

#### Synonyms/Abbreviations

Clostridium difficile PCR, Xpert Clostridium difficile

#### Department

Core Lab

# 2. ANALYTICAL PRINCIPLE

The GeneXpert Dx System automates and integrates sample purification/extraction, nucleic acid amplification, and detection of the target sequence in simple or complex samples using real-time PCR (qPCR) assay. Real-time RT-PCR is used for assays that detect RNA.

The Xpert *C. difficile/Epi* Assay uses real-time PCR to detect DNA. The Xpert *C. difficile/Epi* Assay (where *Epi* means epidemiological) includes reagents for the detection of toxigenic *C. difficile* and the presumptive detection of sequences found in 027/NAP1/BI strains. A Sample Processing Control (SPC) is also included. The SPC is present to control for adequate processing of the target bacteria and to monitor the presence of inhibitors in the PCR reaction. The assay detects the toxin B gene (*tcdB*), the binary toxin gene (CDT), and the single-base-pair deletion at nucleotide 117 within the gene encoding a negative regulator of toxin production (*tcdC* $\Delta$ 117).

# **3. SPECIMEN REQUIREMENTS**

# 3.1 Patient Preparation

Component	Special Notations
Fasting/Special Diets	Not applicable
Specimen Collection and/or Timing	Not applicable
Special Collection Procedures	Transfer liquid or soft stool (but not urine) into the container. Avoid mixing toilet paper, water, or soap with the sample.
Other	None

Criteria		
Type -Preferred	Liquid or semi-formed stool	
-Other Acceptable	None	
<b>Collection Container</b>	Dry sterile leak-proof container	
Volume - Optimum	5 mL	
- Minimum	1 mL	
Transport Container &	Tightly sealed leak-proof container kept	
Temperature		
Stability & Storage	Room Temperature: 24 hours	
Requirements	Refrigerated: 5 days	
	Frozen: Not applicable	
<b>Timing Considerations</b>	Not applicable	
Unacceptable Specimens	• Specimen other than liquid or semi-formed stool	
& Actions to Take	• Specimen with less than 1 mL	
	Specimen past stability requirement	
	• Stool in a wrong transport container	
	• Stool in preservative or mixed with urine	
	<b>Note</b> : Room temperature samples may be tested if received and refrigerated within 24 hours.	
Compromising Physical	Not applicable	
Characteristics		
Other Considerations	Refrigerated samples are to be kept at 2-8°C for up to 5 days.	

# **3.2** Specimen Type & Handling

NOTE: Labeling requirements for all reagents, calibrators and controls include: (1) Open date, (2) Substance name, (3) Lot number, (4) Date of preparation, (5) Expiration date, (6) Initials of tech, and (7) Any special storage instructions. Check all for visible signs of degradation.

#### 4. **REAGENTS**

The package insert for a new lot of kits or reagents must be reviewed for any changes before the kit is used.

#### 4.1 Reagent Summary

Reagents / Kits	Supplier & Catalog Number
Xpert® C. difficile/Epi, GX,	Cepheid, GXCDIFF/EPI-10 (SC#175562) or
IVD Kit	GXCDIFF/EPI-120 (SC#179367) or equivalent

Assay Kit - Xpert® C. diff	Assay Kit - Xpert® C. difficile/Epi, GXCDIFF/EPI-10 or GXCDIFF/EPI-120	
Xpert <i>C. difficile/Epi</i> Assay Cartridges with integrated reaction tubes	Cartridge: • Bead 1 (freeze-dried) • Bead 2 (freeze-dried) • Bead 3 (freeze-dried) • Reagent 1 (3.0 mL per cartridge) • Reagent 2 (3.0 mL per cartridge) – sodium hydroxide	
Xpert <i>C. difficile/Epi</i> Assay Reagent Pouch	1 per kit	
Sample (Elution) Reagent (Guanidinium thiocyanate)	GXCDIFF/EPI-10 x 2.0 mL per pouch GXCDIFF/EPI-120 – 125 x 2.0 mL per pouch	
Storage/Stability	2-28°C / Manufacturer's expiration date Do not use a cartridge that has leaked Do not use a cartridge that has been dropped Do not use a cartridge that has a damaged reaction tube	
Preparation	None required	

# 4.2 Reagent Preparation and Storage

#### 5. CALIBRATORS/STANDARDS

Not applicable

# 6. QUALITY CONTROL

#### 6.1 Controls Used

GeneXpert® C. difficile/Epi PCR Assay	Supplier and Catalog Number
Sample Processing Control (SPC)	Cartridge component
Probe Check (PCC)	Cartridge component
ZeptoMetrix NATtrol <sup>™</sup> <i>Clostridium sordellii</i> External Negative Control	Fisher Cat# 22-156-720; ZeptoMetrix Cat# NATCSO-6MC
ZeptoMetrix NATtrol <sup>™</sup> Clostridium difficile NAP1 External Positive Control	Fisher Cat# 22-156-713; ZeptoMetrix Cat# NATCDI-6MC

# 6.2 Control Preparation and Storage

Sample processing control (SPC) - Included in the Cartridge		
Storage	Refer to section 4	
Stability	Refer to section 4	
Preparation Ready to use		

Probe Check Control (PCC) - Included in the Cartridge	
Container	Refer to section 4
Storage	Refer to section 4
Stability	Ready to use

ZeptoMetrix NATtrol <sup>™</sup> Clostridium difficile NAP1 External Positive Control				
Container	6 x 0.5 mL vials per pack			
Storage	Store at 2–8°C			
Stability	Stable until expiration date.			
Preparation	Control is supplied ready for use. No additional preparation is required.			
	Wearing clean gloves, label 1 cartridge and 1 Elution Buffer appropriately.			
	<ul> <li>Vortex NATtrol<sup>™</sup> control for 5-10 seconds.</li> <li>Add 20 uL NATtrol<sup>™</sup> into Elution Buffer vial.</li> <li>Mix well by vortexing for 10 seconds.</li> <li>Using a sterile transfer pipette, remove all sample from elution buffer and transfer into the "S" chamber of the Assay cartridge. Close cartridge when complete.</li> <li>Control is now ready to be loaded into instrument. Change gloves.</li> </ul>			

#### 6.3 Number and Frequency

- Sample Processing Control (SPC) and a Probe Check Control (PCC; internal controls) are run within each test.
- External *C. difficile* Controls are run with each new kit lot number or shipment or every 31 days, whichever is more frequent. External controls must be treated in the same manner as a patient samples.
- Enter the QC name as QC CDIFF POS and QC CDIFF NEG or scan the QC name barcode

#### 6.4 Tolerance Limits and Criteria for Acceptable QC

A. Tolerance Limits

Control Type	Instrument-Reported Assay Result	Interpretation of Result		
External Positive Control	See Section 10.1	See Section 10.1		
External Negative Control	See Section 10.1	See Section 10.1		
SPC	Passes if Meets the Assigned Acceptance Criteria. Refer to			
PCC	Section 10.1			

- B. Criteria for Acceptable QC
  - All controls must yield acceptable results.
  - Controls and patient data must be reviewed for acceptability and for atypical or unexpected results or trends prior to reporting patient results.
  - DO NOT release results from runs with unacceptable controls or with unusual patterns, trends or distribution in patient values.
- C. Corrective Action
  - Report problem to supervisor or designee.
  - All rejected runs must be effectively addressed and include the following documentation:
    - Control(s) that failed (e.g., positive control with negative result) and/or atypical or unexpected patient results
    - Actions taken
    - Statement of what was done with the patient samples from the affected run/batch,
    - Date and initials of the person recording the information.
  - Patient samples in failed analytical runs must be reanalyzed.

# **NOTE:** The laboratory director or designee may override rejection of partial or complete runs. Justification for the override must be documented in detail.

# 6.5 Documentation

- Record all Quality Control results (failed and successful) manually or electronically.
- Quality control records are reviewed daily at the bench, weekly by the Group Lead or designee, and monthly by the Supervisor/Manager or designee.
- Refer to complete policies and procedures for QC documentation and for record retention requirements in the Laboratory QC Program.
- Refer to Quest Diagnostics Records Management Program for Quality Control record retention requirements.

# 7. EQUIPMENT and SUPPLIES

# 7.1 Assay Platform

• Cepheid GeneXpert System

# 7.2 Equipment

- Computer, monitor, printer, and required application software
- Biological Safety Cabinet
- Timer
- Refrigerator, 2-8°C
- Vortex

• Pipettor – 20uL (for control preparation)

# 7.3 Supplies

- Dry sterile swab
- Sterile transfer pipette
- Aerosol-filter Pipettor tips (for control preparation)
- Plastic-backed absorbent pads (Blood Bloc or equivalent)
- Personal protective equipment (lab coat, powder-free gloves, face shields, and etc)
- Disposable biohazard waste containers (sharps, etc.)
- 10% bleach
- 70% ethanol

# 8. **PROCEDURE**

NOTE: For all procedures involving specimens, buttoned lab coats, gloves, and face protection are required minimum personal protective equipment. Report all accidents to your supervisor.

8.1	Preparation of Cartridge
Notes:	
•	All work must be performed in an appropriate Class 2 BSC (Biological Safety Cabinet).
•	Before testing, clean the work area with a solution of 1:10 dilution of household chlorine bleach and then repeat the cleaning of the work area with 70% ethanol. Wipe work surfaces dry completely before proceeding.
•	Change gloves if they become visually contaminated.
•	Do not open a cartridge until you are ready to perform testing.
•	Use the cartridge within 30 minutes after sample inoculation.
•	Do not use any reagents that have become discolored.
•	Do not touch the integrated reaction tube that is attached to the cartridge.
1.	Remove a test cartridge and Sample Reagent vial from the package and label each with patient specimen number or external control information.
2.	Label the Sample Reagent vial and the Test Cartridge with the accession number.
3.	Briefly place a swab in the liquid/unformed stool sample. The swab does not need to be completely saturated.
4.	Insert the swab into the vial containing the Sample Reagent.
5.	Hold the swab by the stem near the rim of the vial, lift the swab a few millimeters from the bottom of the tube and push the stem against the edge of the vial to break it. Make sure the swab is short enough to allow the cap to close tightly.
	Note: Use clean gauze or plastic-backed absorbent pads for each sample when breaking off swab to minimize risks of contamination.
6.	Replace cap on Sample Reagent and vortex at high speed for 10 seconds.

8.1	Preparation of Cartridge		
7.	Open the cartridge lid. Using a clean transfer pipette, transfer the entire contents of the Sample Reagent to the "S" chamber (labeled 1 below) of the Xpert Assay cartridge.		
8.	Close the cartridge lid and proceed to Section 8.2.		

8.2	GeneXpert Analysis
1.	Turn on the computer, and then turn on the GeneXpert Instrument System.
2.	On the desktop, double-click the GeneXpert software icon.
3.	Log on to the GeneXpert Instrument System software using user name and password.
4.	In the GeneXpert Dx Systems window, click Create Test.
5.	In the Sample ID box, scan or type the accession number (e.g, F1234). Make sure you type the correct sample ID. The sample ID is associated with the test results and is shown in the View Results window and all the reports.
6.	Scan the barcode on the Xpert Assay cartridge.
7.	In the GeneXpert Dx Systems, click Start Test.
8.	Open the instrument module door with the blinking green light and load the cartridge.
9.	Close the door. The test starts and the green light stops blinking. When the test is finished, the light turns off.
10.	Wait until the system releases the door lock before opening the module door and removing the cartridge. Dispose of the used cartridges in a biohazard waste container.
11.	A report is printed for each sample at the completion of testing.

8.3	Retest Procedures			
1.	<ul> <li>If any of the test results mentioned below occur, repeat the test according to the instructions in the Retest Procedures section below.</li> <li>An INVALID result indicates that the SPC failed. The sample was not properly processed or PCR was inhibited.</li> <li>An ERROR result indicates that the Probe Check control failed and the assay was aborted. Possible causes include: the reaction tube being filled improperly; a reagent probe integrity problem was detected; or the maximum pressure limits were exceeded.</li> <li>A NO RESULT indicates that insufficient data were collected. For example, the operator stopped a test that was in progress.</li> </ul>			

8.3	Retest Procedures			
	Retest Procedure			
	For retest within 3 hours of an indeterminate result, use a new cartridge (do not re-use the cartridge) and new reagents.			
2.	<ul> <li>a. Transfer the remaining contents from the Sample Chamber to a new Sample Reagent vial using a disposable transfer pipette.</li> <li>b. Vortex and add the entire contents of the Sample Reagent to the Sample Chamber of the new Xpert <i>C. difficile/Epi</i> Assay cartridge.</li> <li>c. Close the lid and start new test</li> </ul>			
	For retest after 3 hours of an indeterminate result, repeat the test with a new swab sample.			

# **NOTE:** In the event that the test system becomes inoperable, notify supervision or designee for further direction. Patient specimens must be stored in a manner that maintains the integrity of the specimen.

# 9. CALCULATIONS

Not applicable

# 10. REPORTING RESULTS AND REPEAT CRITERIA

#### **10.1** Interpretation of Data

The results are interpreted by the GeneXpert Instrument System from measured fluorescent signals and embedded calculation algorithms and will be shown in the View Results window.

Possible results are:

Result			Internatedian	
Toxin B	<b>Binary Toxin</b>	<i>tcd</i> C	SPC	Interpretation
+	+	+	+/-	Toxigenic C. diff POSITIVE 027-NAP1-BI PRESUMPTIVE POSITIVE
	+	-	+/-	Toxigenic C. diff POSITIVE
+	-	+	+/-	027-NAP1-BI PRESUMPTIVE
	-	-	+/-	NEGATIVE
	+	+	+	Taviania C diffNECATIVE
	+	-	+	Toxigenic C. diff NEGATIVE 027-NAP1-BI PRESUMPTIVE
-	-	+		NEGATIVE
	-	-	+	NEGATIVE

Assay Result Reported	Interpretation of Result			
Toxigenic C. diff POSITIVE; 027 PRESUMPTIVE POSITIVE	<ul> <li>Toxin producing <i>C. difficile</i>, presumptive 027/NAP1/BI target</li> <li>DNA sequences are detected.</li> <li>The toxigenic <i>C. difficile</i> target (Toxin B) AND both presumptive 027/NAP1/BI targets (Binary Toxin and <i>tcd</i>CΔ117) have Cts within the valid range and endpoints above the minimum setting.</li> <li>SPC – N/A; SPC is ignored since <i>C. difficile</i> target amplification may compete with this control.</li> <li>Probe Check – PASS; all probe check results pass.</li> </ul>			
Toxigenic <i>C. diff</i> POSITIVE; 027 PRESUMPTIVE NEGATIVE	<ul> <li>Toxin producing <i>C. difficile</i> target DNA sequences are detected.</li> <li>The toxigenic <i>C. difficile</i> target (Toxin B) AND only one or none of the presumptive 027/NAP1/BI targets (Binary Toxin and <i>tcd</i>CΔ117) have Cts within the valid range and endpoints above the minimum setting.</li> <li>SPC – N/A; SPC is ignored since <i>C. difficile</i> target amplification may compete with this control.</li> <li>Probe Check – PASS; all probe check results pass.</li> </ul>			
Toxigenic C. diff NEGATIVE; 027 PRESUMPTIVE NEGATIVE	<ul> <li>Toxin producing <i>C. difficile</i> target DNA sequences are not detected.</li> <li>Toxigenic <i>C. difficile</i> target (Toxin B) is not detected (regardless of whether Binary Toxin and/or <i>tcd</i>C∆117 is detected).</li> <li>SPC – PASS; SPC has a Ct within the valid range and endpoint above the endpoint minimum setting.</li> <li>Probe Check – PASS; all probe check results pass.</li> </ul>			
INVALID	<ul> <li>Presence or absence of <i>C. difficile</i> target DNA cannot be determined. Repeat test.</li> <li>SPC – FAIL; SPC target result is negative and the SPC Ct is not within valid range and endpoint below minimum setting.</li> <li>Probe Check – PASS; all probe check results pass.</li> </ul>			
ERROR	<ul> <li>Presence or absence of <i>C. difficile</i> target DNA cannot be determined. Repeat test.</li> <li>Toxin producing <i>C. difficile</i> targets — NO RESULT.</li> <li>Binary Toxin (CDT) — NO RESULT.</li> <li><i>tcd</i>CΔ117 — NO RESULT.</li> <li>Probe Check — FAIL*; one or more of the probe check results fail.</li> <li>*If the probe check passed, the error is caused by the maximum pressure limit exceeding the acceptable range.</li> </ul>			

Assay Result Reported	Interpretation of Result	
NO RESULT	<ul> <li>Presence or absence of <i>C. difficile</i> target DNA cannot be determined. Repeat test.</li> <li>Toxin producing <i>C. difficile</i> targets — NO RESULT.</li> <li>Binary Toxin (CDT) — NO RESULT.</li> <li><i>tcd</i>CΔ117 — NO RESULT.</li> <li>Probe Check — N/A</li> </ul>	

# 10.2 Rounding

Not applicable

#### 10.3 Units of Measure

Not applicable

#### **10.4** Analytical Measurement Range (AMR)

Not applicable

#### **10.5** Review Patient Data

- Review patient results for unusual patterns, trends or distribution.
- Report atypical or unexpected results or trends for this test to appropriate supervisory personnel, prior to releasing results.

#### 10.6 Repeat Criteria and Resulting

Repeat Criteria and Resulting			
THEN			
Repeat testing			
Report CDBG as "Detected";			
Add comment PHPV			
Report CDBG as "Detected";			
Add comment NHPV			
Report CDBG as "Not			
Detected"			
Report as INVLD;			
Add comment MPSP			
Г			

Message	Code
Detected	DET
Not Detected	NTD
In addition, the toxigenic C. difficile is PRESUMPTIVELY	PHPV
POSITIVE for a genetic marker of the hypervirulent 027	

Message	Code
NAP1 BI strain, which has been associated with increased toxin production and antimicrobial resistance.	
Simultaneous testing does not identify a genetic marker of the hypervirulent 027 NAP1 BI strain for toxigenic C. difficile	NHPV
After repeat analysis, non-amplification of the internal control suggests the presence of PCR inhibitors in the patient sample. An additional sample should be submitted for testing if clinically warranted.	MPSP
<ul> <li>If C difficile Toxin B is "Detected" then C Diff Comment results as:</li> <li>C difficile toxin and GDH test has been added</li> <li>If C difficile Toxin B is "Not Detected" then C diff Comment is resulted as:</li> <li>C difficile toxin and GDH test not indicated</li> </ul>	*Comment added automatically if <i>C</i> . <i>difficile</i> Toxin B PCR

#### 10.7 Reflex Testing

If the C difficile Toxin B gene (CDBG) is "Detected" then

- Results are held in Sunquest. Call results.
- Quest test C difficile Toxin and GDH(XCDTG) is reflexed. Reflexed test is automatically added to the same accession number as the CDPCR test.
- Reprint XCDTG receipt label or request that accessioning reprint. Label specimen and deliver to accessioning. \*\*when sending specimen to Quest it MUST be **FROZEN**.

# **11. EXPECTED VALUES**

#### **11.1 Reference Ranges**

Not detected

# 11.2 Critical Value

Detected

#### 11.3 Standard Required Messages

None established

# **12.** CLINICAL SIGNIFICANCE

Clostridium difficile (C. difficile) is a Gram-positive, spore-forming anaerobic bacillus that was first linked to disease in 1978. C. difficile infection (CDI) ranges from diarrhea to severe life-threatening pseudomembranous colitis. C. difficile's primary virulence factor is cytotoxin B. The genes coding for toxin A (tcdA; the enterotoxin) and toxin B (tcdB) are parts of the pathogenicity locus (PaLoc). Most pathogenic strains are toxin A-positive, toxin B-positive (A+B+) strains although toxin A-negative, toxin B-positive (A-B+) variant isolates have been recognized as pathogenic. Some strains of C. difficile also produce an actin-specific ADP-ribosyltransferase called CDT or binary toxin. The binary toxin locus contains two genes (cdtA and cdtB) and is located outside the PaLoc.

In the last several years, there have been outbreaks of CDI attributed to a number of emerging "hypervirulent" strains that include fluoroquinolone resistant strains belonging to PCR ribotype 027, PFGE type NAP1 and REA type BI. Strains of 027/NAP1/BI exhibit increased toxin production, which is being attributed to deletions in the regulatory gene *tcdC* and they are thought to produce more spores, leading to enhanced persistence in the environment. The identification of a presumptive positive or negative 027/NAP1/BI result may aid in the identification of possible sources of an 027/NAP1/BI outbreak.

*C. difficile* diagnosis has been traditionally based on the detection of toxin A or B. Both the labor intensive culture procedure, followed by cell cytotoxicity testing on the isolates, and cytotoxicity cell assay on stool specimens are still considered to be the "gold standard" because of high specificity. Several rapid enzyme immunoassays have been developed for detection of toxin A and B. However, these tests have reduced sensitivity and specificity compared to the cell cytotoxicity assay. Recently, PCR methods for the detection of toxin A and/or toxin B have been developed with high sensitivity and specificity as compared to the cell cytotoxicity and immunoassays.

# **13. PROCEDURE NOTES**

- FDA Status: FDA Exempt/Cleared or Approved
- Treat all biological specimens, including used cartridges, as if capable of transmitting infectious agents. Because it is often impossible to know which might be infectious, all biological specimens should be treated with standard precautions.
- Follow your institution's safety procedures for working with chemicals and handling biological samples.
- In the event of contamination of the work area or equipment with samples or controls, thoroughly clean the contaminated area with a solution of 1:10 dilution of household chlorine bleach and then repeat the cleaning of the work area with 70% ethanol. Wipe work surfaces dry completely before proceeding.
- Results from Xpert *C. difficile/Epi* Assays are NOT intended to guide treatment of *C. difficile* infections.
- Performance characteristics were not established for patients < 2 years of age.

- The Xpert *C. difficile/Epi* Assay does not provide susceptibility results. A separate specimen aliquot and additional time are required to culture and perform susceptibility testing.
- Do not substitute Xpert C. difficile/Epi Assay reagents with other reagents.
- Do not open the Xpert *C. difficile/Epi* Assay cartridge lid except when adding sample and reagents or performing a retest.
- Do not use a cartridge that has been dropped.
- Do not use a cartridge that has a damaged reaction tube.
- Each single-use Xpert *C. difficile/Epi* Assay cartridge is used to process one test. Do not reuse spent cartridges.

# 14. LIMITATIONS OF METHOD

#### 14.1 Precision

Not applicable

#### 14.2 Interfering Substances

As indicated in the package insert, twenty-one (21) biological and chemical substances occasionally used or found in stool specimens were tested for interference with the Xpert *C. difficile/Epi* Assay. Potentially interfering substances include, but are not limited to Vagisil cream and zinc oxide paste (see "Assay Limitations"). The 19 substances listed below showed no detectable interference with the Xpert *C. difficile/Epi* Assay.

Substance	Substance
Whole Blood	K-Y Jelly/Gelée
Mucin (porcine)	Vaseline
Kaopectate	Dulcolax
Imodium	Preparation H Portable Wipes
Pepto-Bismol	Vaginal Contraceptive Film (VCF)
Preparation H	Vancomycin
Fleet	Metronidazole
Fecal fats	Anusol Plus
Monistat	E-Z-HDTM High Density Barium Sulfate for
	suspension
Hydrocortisone Cream Longs Drugs	

# 14.3 Clinical Sensitivity/Specificity/Predictive Values

As indicated in the Package Insert, the Xpert *C. difficile/Epi* assay had overall sensitivity, specificity, positive predicative value, and negative predicative value of 88.7%, 90.9%, 55.4%, and 99.8% respectively when compared to direct culture with strain typing.

- Non-027/NAP1/BI isolates representing toxinotype XIV will be reported "Toxigenic *C. diff* POSITIVE; 027 PRESUMPTIVE POSITIVE" using the Xpert *C. difficile/Epi* Assay.
- Occasionally, non-027/NAP1/BI isolates representing toxinotypes IV, V and X will be reported "Toxigenic *C. diff* POSITIVE; 027 PRESUMPTIVE POSITIVE" using the Xpert *C. difficile/Epi* Assay.
- The performance of the Xpert *C. difficile/Epi* Assay was validated using the procedures provided in this package insert only. Modifications to these procedures may alter the performance of the test.
- Results from the Xpert *C. difficile/Epi* Assay should be interpreted in conjunction with other laboratory and clinical data available to the clinician.
- 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. Careful compliance with the instructions in this insert is necessary to avoid erroneous results.
- Because of the dilution factor associated with the retest procedure, it is possible that *C. difficile* positive specimens, very near or at the limit of detection (LoD) of the *C. difficile/Epi* Assay, may result in a false negative result upon retest.
- Inhibition of the Xpert *C. difficile/Epi* Assay has been observed in the presence of the following substances: Zinc oxide paste and Vagisil® cream.
- Outbreaks of CDI may be caused by strains other than 027/NAP1/BI.
- False-negative results may occur when the infecting organism has genomic mutations, insertions, deletions, or rearrangements or when performed very early in the course of illness.

# 15. SAFETY

- Reagent 1 contains sodium hydroxide (pH > 12.5); (R34 EU Risk) which is corrosive to eyes and skin requiring eye and skin protection.
- Refer to your local and corporate safety manuals and Safety Data Sheet (SDS) for detailed information on safety practices and procedures and a complete description of hazards.

# **16. RELATED DOCUMENTS**

- Biological Safety Cabinet, Micro procedure
- Laboratory Quality Control Program
- Laboratory Safety Manual
- Safety Data Sheets (SDS)
- Quest Diagnostics Incorporated Records Management Procedure
- *Clostridium difficile* Toxin B PCR using Cepheid® GeneXpert (QDMD734)
- GeneXpert Dx System Operator Manual
- Cepheid GeneXpert® Dx System Maintenance, Micro procedure
- *Clostridium difficile* PCR Quality Control Log (AG.F410)

• Cepheid GeneXpert<sup>®</sup> C. difficile Toxin B PCR Individual Quality Control Plans (SGAH.VC371, WAH.VC253)

# **17. REFERENCES**

- 1. Xpert® MRSA Assay current package insert (11/2012).
- 2. American Society for Microbiology. 2010. A Practical Guidance Document for the Laboratory Detection of Toxigenic *Clostridium difficile*.
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- 6. Bignardi GE. Risk factors for Clostridium difficile infection. J Hosp Infect. 1998; 40:1-15.
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# **18. DOCUMENT HISTORY**

Version	Date	Section	Revision	Revised By	Approved By
			Supersedes SGAHQDMD734v1.3		
1	6/8/20	Header	Changed WAH to WOMC	L Barrett	R Master
		8.2	Deleted typing patient name and MRN into instrument		
		10.6	Added interfaced reporting		
		19	Added addendum A		

Version	Date	Section	Revision	Revised By	Approved By
2	1/11/22	10.6	Removed comment message, when C diff tox Detected and replaced to with new ones, applicable to new reflex test	M Sabonis	R Master
			Removed steps for results from 10.6 and moved to Addendum A		
			Added Sunquest example screen shots for resulting positive C diff results with reflexing		
2	1/11/22	10.7	Added steps for reflex testing.	M Sabonis	R Master
2	3/8/22	Footer	Changed prefix to AHC	D Collier	R Master
2	3/8/22	8.1	Spelled out Biological Safety Cabinet	C Bowman- Gholston	R Master

# **19. ADDENDA**

A. Cepheid Testing and Running via Sunquest Interface

# Addendum A

# Cepheid Testing and Running via Sunquest Interface

# A. General Information:

- 1. This interface does NOT go through DI-Instrument Manager. Cepheid is interfaced directly to Sunquest. The Sunquest interface is set up for Autoverification.
- 2. All tests will auto-file except for those that must be called.
- 3. If the test is positive for *C. difficile*, then the results will be <u>held</u> in Sunquest. These results must be called and documented per routine process.
- 4. Use function OEM on Sunquest SmarTerm to review results via the interface.
  - a. Access OEM
    - At DEVICE: prompt, type in Method code **WOCE** (WOMC) or **SGCE** (SGMC).
    - Results will display cup by cup.
      - $\circ$  Those that were auto-filed require no action, proceed to next cup.
      - For positive results that were held, continue with steps b and c below.
    - Refer to *OEM On Line Result Entry Method* procedure (LIS SOP) for additional information about review and release of results.
  - b. Call results. Append CBACK documentation to results including who you called, date, time and tech code. Required format is:

-CBACK-;full name of person called DATE TIME Tech code *Example* -CBACK-;Sue Smith 032420 1420 4568

c. Click on Accept to release results.

# Below is example from SmarTerm OEM, when C difficile results is "Detected"

DEVICE LOC: WAH WASHINGTON AD	DNLINE RESULT ENTRY DVENTIST HOSPITAL	HOSP. ID: WAH
CUP 581 ACC NO NAME PN: 1	TEST-50 AGE/SE	X LOC PHYSICIAN
W2995 TEST,MARIE		TEST CACCIABEVE,NICO
	DOB: 06/26/2018	COLL: 01/12/2022 06:08
F	DETECTED FAILED NORMAL [NTD] Call Results	Message displays: *XCDTG reflexed, send to Quest *Call results
(ADD) TEST-1: XCDTG-OBL	ordered & adde	CDTG is being ed to the same
Orders for dept: General Lab Test(s): CDPCR XCDTG-OBL	accession #	
ACC. NO: W2995	Press RETUR	RN to continue
ACC. NO: W2995 REQUEST COMPLETE RETURNING	TO ONLINE RESULT ENTR	۲Y
CDHV : NHPV	ic marker of the hype ain for toxigenic C.d	
CDCMT : CDTA	C difficile toxin and	GDH test has been added
(ORDER: CDPCR)	testin	sage posts denoting reflex ng has been added
ACCEPT (A), MODIFY (M), DISPL	AY PRIOR (D), PRE	

- 5. Perform an OFC (Online File Cleanup) at least once per shift. This process cleans up the online data that was sent to Sunquest.
  - a. In Sunquest (SmarTerm) access function OFC
  - b. Type in the method code (WOCE or SGCE).
  - c. At the Start at Cup Number prompt, type in 1 and then press ENTER.
  - d. At the Stop at Cup Number prompt, press ENTER.
- 6. Use function MEM, if manually entering results into Sunquest
  - a. SmarTerm: function **MEM** to enter results.
  - b. Enter Shift (1, 2, or 3)
  - c. Worksheet: Use WIM2 for WOMC or SIM2 for SGMC.
  - d. Test: <Enter>
  - e. Enter "A" (Accept)
  - f. Enter Accession number
  - g. Press <Enter> until Result screen displayed
  - h. Key in result using appropriate code from above

# **B.** Running Tests on Cepheid:

- 1. Create Test
  - a. In the GeneXpert Dx System window, click **Create Test** on the menu bar. The Scan Sample ID Barcode dialog box appears.

per Data Manager	ment Re	eports Setup Maintenance Abor				20		User lab
ASA			0	()		000	La	H
Create Tes		Check Status	Stop Test	View Results	Define	Assays	Define Graptis	Maintenance
		Modules	1	In order the design of the		Tests Since Launch		
Module An	150Y	Create Test	Transmiss.	inensie seer	18985	meast doors		r Start , Dute
Hamo Al		Host Test Order Table	The second second				And the second second second	01/22/20 09:27:48
A1 A2		Sample ID	Assay	STAT	Host Ord	ter Time		01/22/20 07:58:24
A3								01/22/20 07:47:47
A4		and the second second						01/22/20 07:19:31
81								01/22/20 06:53:56
82				Galete Ak Host Test				01/21/20 21:45:40
83	1000			and the second of the second second	Concession and the second second			01/21/20 17:33.28
84							Manual Query	01/21/20 16:35:46
	1.00	1				*		01/21/20 16:32:30
		The second states of the second	6.0	mole ID Barcode	X			01/21/20 15:54:08
			Scan Sa	mple ID sarcode	Contraction of Contraction			01/21/20 13:36:06
		Sample ID						01/21/20 10:19:20
				scan sample ID barcode.		The Party of the		01/21/20 09:43:02
	101		Name					01/21/20 01:02:04
	100	Select Assay	-None>				-	01/20/20 23:24:26
	1810	Select Module				and the second second		01/20/20 21:07:54
	3143			Monual Entry	Cancel		The second s	01/20/20 19:29:48
		Reegent Lot ID						04000046-3046
	C ST LAND	Test Type	Specimen		Concess Manual make	and the second second		
dule A1: Lest Star	ned at UT		abacautan					
ase load the carb	vidge Inte	Sample Type	I	<b>•</b> 08	her Sample Type			
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dule A3: Test Star	rted at 01						and the second	
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odule B2: Test Stat ease load the carb	integ at 01			Scan Cartridge Ba	arcode Cancel			
odule 83: Test Stat	rted at 01	Lange de la companya de la					Contract of the second	9

b. Scan the Sunquest barcode label.

GeneXpert® Dx Syste					_ 0 ×
User Data Management F	Reports Setup Maintenance About	A training of the second s			User labsuper
A CONTRACT	M	Ø		Define Graphs	Madeauxe
Create Test	Check Status Modules	Slop Test View F	Kesuite Define Assayo Tests Sim		to AUTOCALCE
Module Assey A1 A2	Create Test Host Test Order Table Sample ID		STAT Host Order Time	X	Indicates the Cepheid
A3 A4 B1 82	040000775 Xpert C diff	Epi Version 2 Normal	01/22/20 10:42:08	Detete	queried Sunquest and found
B3 B4	Host query for sample (040300775) sent at 0 1 order(s) have been downloaded. Query completes at 01/22/20 10:42:08.			Manael Query	the order. In this example the order was for a C
	Sample ID <sup>4</sup> 04000077 Name Salact Assay	Scan Cartridge Barcode			difficile PCR (Xpert C diff- Epi)
A <del>.</del>	Select litodule A2 Reegent Lot ID	Manual Entry	Cancel		01720/2019/23/48
Messages: Module AT: Lest Stated at 0 Please load the cartridge into Module A2: Test Stated at 0 Please load the cartridge into Module A3: Test Stated at 0 Please load the cartridge into Module A4: Test Stated at 0 Please load the cartridge into	Test Type Spectma Sample Type Other Hotes	• •	Other Sample Type		
Module B1: Test Started at 0 Please load the cartridge into Module B2: Test Started at 0 Please load the cartridge into Module B3: Test Started at 0 Please load the cartridge into		Start Test Scan Cartrid	ige Barcode Cancel		

- c. Scan the cartridge barcode.
- 2. Click OK
- 3. Click Create Test
- 4. Load cartridge
- 5. Verify that the test has started before walking away
- 6. When testing is completed results will print to Cepheid printer.

#### C. Manually uploading results to Sunquest (Example Sunquest downtime)

- 1. From the Cepheid, go to VIEW RESULTS
  - a. Click on UPLOAD TEST and find the Sample ID (Sunquest Accession #).

	Sample (D*		Views		Analyte Result Detail Errors History Support
030030922			Result View Primary Curve		Xpert MRSA NxG Version 2
Assay Version Test Type Sample Type	Specimen	• •		Test Result	IIII SA NOT GETTETEL
Othe	r Sample Type			For In Vitro Dia	sgnostic Use
	Notes	100			
8255512, PAPAN	ICOLAS		Views		Legend
Upload Status Module Name Reagent Lot ID*	B2		Result View Primary Curve	60 8	IC IC IC IC IC IC IC IC IC IC
End Time Status	01/29/20 01:05:40 01/29/20 02:14:27 Done labsuper			40 Hundescence	
					0 10 20 30 40 Cycles
Save Charges	Export	port	Upload Test	Seguri Graphy	View Test

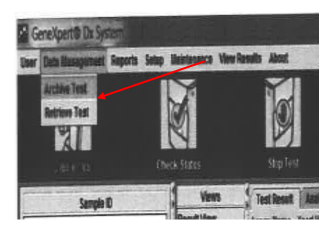
- b. Check off the one that you want to upload (located to the left of the Update Status column). Note: You can check off one or more accession numbers at the same time.
- c. Click on **UPLOAD** to resend to Sunquest. Results will now upload into Sunquest. It make take a little time for upload to complete.

	Upload Status	STAT	Semple	User	Result	Assay	Status	Error	Start
	Uploaded	Normal	050050850	labsuper	MRSA NOT DETE	Xpert MRSA NxG	Done	OK	01/30/20 08:20:13
	Uploaded	Normal	040033476	labsuper	MRSA NOT DETE	Xpert MRSA NxG	Done	OK	01/30/20 07:57:52
	Uploaded	Normal	050051032	labsuper	MRSA NOT DETE	Xpert MRSA NxG	Done	OK	01/30/20 07:48:14
	Uploaded	Normal	050050907	labsuper	MRSA NOT DETE	Xpert MRSA NxG	Done	OK	01/30/20 07:25:27
	Uploaded	Normal	050050626	labsuper	MRSA NOT DETE	Xpert MRSA NxG	Done	OK	01/30/20 06:25:16
	Uploaded	Normal	050050876	labsuper	MRSA NOT DETE	Xpert MRSA NxG	Done	OK	01/30/20 06:24:14
	Uploaded	Normal	H-0288	labsuper	MRSA NOT DETE	Xpert MRSA NxG	Done	OK	01/30/20 04:30:00
	Uploaded	Normal	H50 32	labsuper	MRSA NOT DETE	Xpert MRSA NxG	Done	OK	01/30/20 04:28:37
	Uploaded	Normal	050050358	labsuper	MR SA MISSA NOT D	ANXG	Done	OK	01/30/20 03 55:09
	Uploaded	Normal	W34777	labsuper	Toxigenic C.diff P		Done	OK	01/30/20 01 56:48
	Uploaded	Normal	04003254	labsuper	MRSA NOT DETE	Xpert MRSA NxG	Done	OK	01/30/20 00:55:21
	Uploaded	Normal	050050307	labsuper	MRSA NOT DETE	Xpert MRSA NxG	Done	OK	01/30/20 00:53:39
	Uploaded	Normal	040034494	labsuper	MRSA NOT DETE	Xpert MRSA NxG	Done	OK	01/29/20 22:07:15
	Uploaded	Normal	-W33590	absuper	MRSA NOT DETE	Xpert MRSA NxG	Done	ок	01/29/20 19:05:10
	Uploaded	Normal	040033585	labsuper	MRSANOT DETE	Xpert MRSA NxG	Done	OK	01/29/20 17:55:21
	Uploaded	Normal	040033524	labuper	MRSA NOT DETE	Xpert MRSA NtG	Done	OK	01/29/20 16:23:38
	Uploaded	Normal	040033177	labsuper	MRSA NOT DETE	Xpart MRSA NxG	Done	OK	01/29/20 14:27:52
	Uploaded	Normal	040032800	labsupur	MRSA DETECTED	Xpert MRSA NxG	Done	OK	01/29/20 12 29:06
Ē		, r					-	1 1	*******
	Select		Deselect All		Select Highlighted		Deselect Highlighted	1000	Select All Pending

d. Review in Sunquest OEM to document any positive result call notification.

#### D. Editing Sample ID (SQ Accession #)

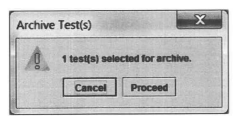
1. From the main screen - $\rightarrow$ Data Management- $\rightarrow$  Click on Archive Test



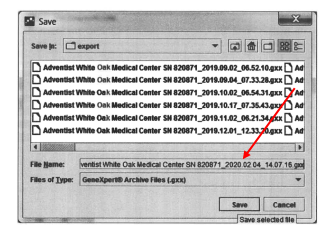
2. In the upper left corner click on **Purge Selected Tests from the LIS after Archiving** (red arrow). Then locate the Sample ID (SQ Accession#) that you want and select it by clicking on box to the left of the Sample ID (blue arrow). Then click on OK.

-	Sample	T Module	User	Resut	Азвау	States	Error Status	Start Date
	M19573	83	labsuper	MRS'OT DETECTED	Xpert MRSA NxG	Done	OK	01/08/20 23:03:45
<u>–</u>	M19573	84	labsuper	RSA NOT DETECTED	Xpert MRSA NxG	Done	OK	01/06/20 23:38:24
	H50632	A1	labsuper	MRBA NOT DETECTED	Xpert MRSA NxG	Done	ок	01/30/20 04:28:37
Contraction of the	H50288	A2	labsup	MRSA NOT DETECTED	Xpert MRSA NxG	Done	OK	01/30/20 04:30:00
	H45323	A1	19dtper	MRSA NOT DETECTED	Xpert MRSA NxG	Done	OK	01/17/20 04:14:48
<u> </u>	F36575	83	labsuper	MRSA NOT DETECTED	Xpert MRSA NxG	Done	OK	01/31/20 23:38:17
	F33568	A3	labsuper	MRSA NOT DETECTED	Xpert MRSA NxG	Done	OK	01/31/20 05:03:19
- <u>H</u>	F33294	15	labsuper	MRSA DETECTED	Xpert MRSA NxG	Done	OK	01/31/20 05:12:25
-H	F28121	150	labsuper	MRSA NOT DETECTED	Xpert NRSA NxG	Done	OK	01/18/20 00:10:12
	F20121	12	labsuper	MRSA NOT DETECTED	Xpert MRSA NxG	Done	OK	01/18/20 02:42:55
<u> </u>	F27167	RA BA	labsuper	MRSA NOT DETECTED	Xpert MRSA NxG	Done	OK	01/17/20 04:12:18
H	F20193	83	labsuper	NO RESULT	Xpert MRSA NtG	Stopped	OK	01/17/20 04:08:12
<b>e</b>	F25041	81	labeuper	MRSA NOT DETECTED	Xpert GRSA NoG	Done	IOK	01/17/20-01:44:50
	46762557	84	labsuper	Toxigenic C.diff POSITIV.	Xpert C.dlf-Epi	Done	OK	01/20/20 07:12:56
- <u>H</u>	070016548	A2	labsuper	MRSA NOT DETECTED	Xpert MRSA NxG	Done	OK	02/01/20 22:27:33
	070010040							
-	Select		eselect	Select		Deselect		Select New

3. At the Archive Test prompt, click on **Proceed**.



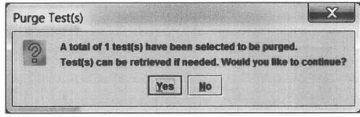
4. Archive file is generated (File name is system generated) and click on **SAVE**. Note that the File Name has the date and time as part of the file name. In the example below "2020.02.04\_1407" is the date of 2/4/20 and time of 1407.



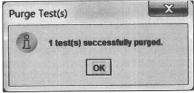
5. Archive message displays, click on OK

1 test(s) successfully archived and can be found at     C::GeneXpertlexportAdventist White Oak Medical Center SN 820871_2020.02.04_14.07     OK	X
OK	07.16.gxx.

6. Purge message displays, click on OK



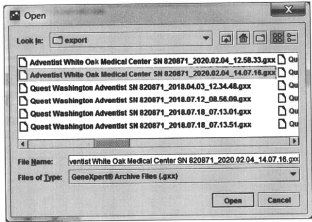
7. Completion of purge message displays



8. Retrieve test by going to Main screen - $\rightarrow$  Data Management- $\rightarrow$  Retrieve Test



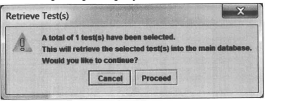
9. Locate file that you exported (Note, part of the file name consists of the date and time file was created.). Highlight the file and click on Open.



10. To the left of the Sample ID, check off the Sample ID (SQ acc #) that you want to retrieve to edit. Then click on OK.

	Sample		Module Name	User	Result	Assay	Stains	Error Status	Start Date
8	F25041	B1		labsuper	MRSA NOT DETECTED	Xpert MRSA NxG	Done	ок	01/17/20 01:44:50
	Sellinct		Dese		Select		Deselect		Select With
	Select All		Dese		Select Highlighted		Deselect Highlighted	[	Select With No Duplicate

11. Retrieve prompt displays. Click on Proceed. Retrieve Test(s) confirm displays. Click on OK.

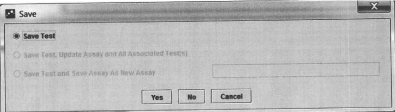


Retrieve	e Test(s)		
ï	1 test(s) successfully retrieved. Retrieval details are logged in C:(GeneXpert(Database_log)Retrieve_log_2020.02.04_14.12.46.bx		
	OK		

12. Proceed to edit Sample ID (SQ Accession #). Click on Save when you are done.

ter Dets Man	gement Reports 5	Check
Create	Sample ID	
F25041		
Assay Version	Xpert MRSA NxG	
Test Type	Specimen	-
Sample Type	Other	*
Othe	er Sample Type	
Refer thanks	Notes	
STEWART,G 176364		
Module Name	B1	
Reagent Lot 10*	05306	
11 1 - ALTERNATOR STATISTICS	01/17/20 01:44:50	
End Time	01/17/20 02:53:52	1.12
	Done .	22
User	labsuper	
1.10		

13. Click on **Yes** on the Save Test prompt.



14. Follow the steps in part C above to upload the results to Sunquest.