

Assay Technical Training: Xpert[®] CT/NG For CE-IVD and US-IVD Use





In Vitro Diagnostic Medical Device

301-1883 Rev. F, August 2019

Training Agenda

Ē

Xpert[®] CT/NG Training

- Intended use
- Reagents
- Sample collection
- Kit storage and handling
- Preparing the cartridge
- Quality Controls
- Results Analysis
- Discussion





Training Objectives

- At the end of the training, users will be able to:
 - Store and handle the Xpert[®] CT/NG cartridge kit and Sample collection kits
 - Follow proper laboratory safety precautions
 - Collect appropriate specimen types and transport specimens
 - Perform the cartridge set up and run the assay
 - Report the various software-generated results
 - Understand the assay control strategy



The Cepheid Solution



- Simultaneous detection of CT and a dual target NG (NG2/NG4)
- On-board controls for each individual sample
 - Probe Check Control (PCC)
 - Specimen Processing Control (SPC)
 - Sample Adequacy Control (SAC)
- Results in approximately 90 minutes
- Closed cartridge system minimizes risk of contamination
- On-demand results
- Random access





Intended Use

The Xpert [®] CT/NG Assay is an automated in vitro diagnostic test for qualitative detection and differentiation of DNA from *Chlamydia trachomatis* (CT) and *Neisseria gonorrhoeae* (NG).

The assay may be used to test the following specimens from asymptomatic and symptomatic individuals:

- Specimen
 - Urine Specimen (male and female)
 - Pharyngeal Swab Specimen (male and female)
 - Rectal Specimen (male and female)
- Detects:
 - CT target sequence
 - NG (NG2/NG4) target sequences

- Patient-collected Vaginal Specimen (collected in a clinical setting)
- Endocervical Specimen





Targets

5 targets are detected:

- CT1 located on genomic DNA (also present in the genome of the swedish variant strains of Chlamydia trachomatis)
- NG2 independent and unique target of Neisseria gonorrhoeae
- NG4 independent and unique target of Neisseria gonorrhoeae
- SPC
- SAC



Assay Requirements

GeneXpert[®] Systems

• GeneXpert[®] Dx Software v4.3 or higher/ Xpertise[™] Software v6.0 or higher

Test Kits

- GXCT/NG-10 and GXCT/NG-120 (US-IVD)
- GXCT/NGX-CE-10 and GXCT/NGX-CE-120 (CE-IVD)

Sample Collection

- SWAB/A-50
 - SWAB/G-50
- URINE/A-50

Other materials

- Personal Protective Equipment (PPE)
- 1:10 dilution of household bleach
- 70% ethanol or denatured ethanol

Optional

- Uninterruptible Power Supply /Surge Protector
- Printer



Good Laboratory Practice



Kit Handling



Xpert[®] CT/NG Assay Kit Contents

Catalog Number	GXCT/NG-10 or GXCT/NGX-CE-10 GXCT/NG-120 or GXCT/NGX-CE-120		
Tests Per Kit	10 or 120		
Kit CD	Assay Definition File (ADF)		
	Assay Import Instructions		
	Package Insert (PDF)		
Transfer Pipettes	10 or 120		
Storage	2-28 °C		

Note: Sample Reagent contains guanidinium thiocyanate, which is harmful if swallowed (H303) and Irritating to eyes and skin (H315, H319).

Cartridges contain chemically hazardous substances-please see Package Insert and Safety Data Sheet for more detailed information.







Warnings and Precautions

Store the Xpert[®] CT/NG Assay cartridges and reagents at 2–28°C



- Follow your institution's safety procedures for working with chemicals and handling biological samples
- Do not use Collection Reagent tubes that have not been validated by Cepheid
- Open the Assay cartridge lid only when adding the Sample, close the lid and proceed with processing
- With the GeneXpert System, start the test within 30 minutes after adding the sample to the cartridge
- With the Infinity System, place the cartridge on the conveyor within 30 minutes of adding the sample.



Warnings and Precautions

- Do not shake the cartridge
- Do not use a cartridge that...:
- appears wet, has leaked or if the lid seal appears to have been broken
- appears damaged
- has been dropped after removing it from packaging
- has been dropped or shaken after adding the sample to it
- has a damaged reaction tube
- has been used: each cartridge is single-use to process one test
- is expired
- do not reuse disposable pipettes

Dispose Xpert* CT/NG Assay cartridges and reagents according to your institution's and country's guidelines for disposal of hazardous materials

Warnings and Precautions

• Biological specimens, transfer devices, and used cartridges should be considered capable of transmitting infectious agents requiring standard precautions.

• Follow your institution's environmental waste procedures for proper disposal of used cartridges and unused reagents. These materials may exhibit characteristics of chemical hazardous waste requiring specific national or regional disposal procedures.

• If national or regional regulations do not provide clear direction on proper disposal, biological specimens and used cartridges should be disposed per WHO [World Health Organization] medical waste handling and disposal guidelines.



Ē

Xpert® CT/NG Assay Limitations

- The Xpert CT/NG test has been validated with the following specimen types, collected with the Xpert Swab Specimen Collection Kit, Xpert CT/NG Urine Specimen Collection Kit or Xpert Urine Specimen Collection Kit.
 - Endocervical swabs
 - Patient-collected vaginal swabs
 - Male and female pharyngeal swabs
 - Male and female rectal swabs
 - Male and female urine
- Presence of vaginal discharge, tampons, douching, and Cepheid non-validated specimen collection have not been determined.
- Collection and testing of urine specimens with the Xpert[®] CT/NG Assay is not intended to replace a cervical exam and endocervical sampling for diagnosis of urogenital infections. Other genitourinary tract infections can be caused by other infectious agents.
- The performance of Xpert[®] CT/NG has not been evaluated for patients less than 14 years of age or in patients with a history of hysterectomy.
 CE-IVD and US-IVD. For *in vitro* diagnostic use
 For detailed information, refer to the current Package Insert

Specimen Collection, Storage and Transport

Cepheid

Urine Specimen Collection

• Urine

- Refer to urine specimen collection package insert
- 20 to 50 ml of first-catch urine should be collected in a sterile urine collection cup with no preservative (not provided by Cepheid), from which 7 mL is transferred to the Urine Sample tube containing the preservative
- Use only Xpert Urine Sample Collection kit for processing male and female urine prior to testing in the GeneXpert CT/NG assay
- Urine specimen must be collected and tested before the expiration date of the Xpert Urine Sample Collection kit







*Cepheid catalog # URINE/A-50

Urine Specimen Collection (First Catch)



Direct patient to provide first catch urine (20-50 mL) into a urine collection cup. Note: The patient should not have urinated for at least 1 hour before. Patient should not cleanse the genital area prior to collecting specimen.





Open the package of disposable transfer pipette.



Remove the yellow cap from the transportation tube.



Transfer approximately 7 mL of urine into the transport tube, using the disposable transfer pipette. The correct volume is marked by the black dashed line on the pipette.



Replace the yellow cap on the transport tube and tighten securely.



Invert the transport tube 3-4 times to ensure that the specimen and reagent are well mixed.



Return the tube as instructed by your doctor, nurse or health care provider.



Under or over dispensing of urine into the Xpert Urine Transport Reagent tubes may affect assay performance

Specimen Collection, Transport and Storage

Unprocessed Urine samples

Ē





Specimen Collection, Transport and Storage

Urine samples transferred to Xpert[®] CT/NG Urine transport tube

Cepheid catalog # URINE/A-50

Ē

			Temperature (°C)	Storage Time
	Female urine sample	Ç	+15 °C	45 days
	Female urine sample	Ç	+ <u>2</u> + <u>30</u> °C	3 days
	Male urine sample	ď	+ <u>2</u> + <u>30</u> °C	45 days



Vaginal and Endocervical Collection

• Swab

- Pharyngeal, rectal, vaginal and endocervical specimens are collected from patients using flocked swabs included in the kit
- Vaginal samples are collected by the patient.
 Conversely, endocervical samples are collected by a clinician
- Swabs are broken off into the transport reagent tubes to elute organisms and stabilize DNA
- Swab specimens are then transported to the laboratory for testing on the GeneXpert[®] Instrument

Volume of transport medium: 2,4 mL



*Cepheid catalog # SWAB/A-50 for vaginal & endocervical samples

or Cepheid catalog # SWAB/G-50 kits for pharyngeal, rectal, vaginal & endocervical sample

SWAB/A-50 and SWAB/G-50 kits are designed to collect, preserve and transport endocervical and vaginal specimens from symptomatic and asymptomatic individuals to the lab prior to analysis with the Xpert[®] CT/NG Assay



Pharyngeal Collection



Open the individual rectal specimen collection package that contains the pink-capped swab transport tube and individually wrapped collection swab. Discard the larger swab



Open the collection swab wrapper by peeling the top of the wrapper



Hold the swab in your hand, placing your thumb and forefinger in the middle of the swab shaft across the scoreline.



Re-cap the transport tube and tighten the cap securely.



Instruct the patient to open mouth widely. Position the tongue toward the bottom of the mouth. Swab areas of the pharynx (tonsil, posterior wall, uvula, posterior wall).

Invert or gently shake the tube 3-4 times to elute material from the swab. Avoid foaming. Label the transport tube with the sample ID, including date of the collection, as required.

Avoid splashing contents of the transport tube on the skin. Wash with soap and water if exposed.





While holding the swab in the same hand, unscrew the cap from the Xpert Swab Transport Reagent tube.



Carefully break the swab shaft against the side of the tube at the scoreline

Rectal Collection



Open the individual rectal specimen collection package that contains the pink-capped swab transport tube and individually wrapped collection swab. Discard the larger swab



Open the collection swab wrapper by peeling the top of the wrapper



Hold the swab in your hand, placing your thumb and forefinger in the middle of the swab shaft across the scoreline.



Carefully insert the swab approximately 1 cm beyond the anal sphincter, so that the fiber tips are no longer visible and rotate the swab.



While holding the swab in the same hand, unscrew the cap from the Xpert Swab Transport Reagent tube.



Carefully break the swab shaft against the side of the tube at the scoreline



Re-cap the transport tube and tighten the cap securely.

Invert or gently shake the tube 3-4 times to elute material from the swab. Avoid foaming. Label the transport tube with the sample ID, including date of the collection, as required.

Avoid splashing contents of the transport tube on the skin. Wash with soap and water if exposed.



Assessing a correct sample

Figure 1. Examples of Acceptable Rectal Swabs



Figure 2. Examples of Unacceptable Rectal Swabs





Vaginal Collection (Patient Collected)



Open the individual vaginal/endocervical specimen collection package Athat contains the pink-capped swab transport tube and individually wrapped collection swab. Set the tube aside. Discard the larger B swab



Open the collection swab by peeling open the top of the wrapper.

Remove the swab, taking care not to touch the tip or lay it down.



Hold the swab in your hand, placing your thumb and forefinger in the middle of the swab shaft across the scoreline.



Carefully insert the swab into your vagina about 2 inches/5cm inside the opening of the vagina



Gently rotate the swab for 10-30 seconds. Ensure the swab touches the walls of the vagina so that the moisture is absorbed by the swab.

Withdraw the swab and continue to hold it in your hand.



Unscrew the cap from the transport tube. Immediately place the collection swab into the transport tube.



Identifying the scoreline, break the swab shaft against the side of the tube. Discard the top portion of the swab shaft. Avoid splashing contents on the skin. Wash with soap and water if exposed.



Re-cap the transport tube and tighten the cap securely. Return the tube as instructed by your doctor, nurse or health care provider.



Endocervical Collection (Clinician-collected)



Cleaning Swab

B



Remove excess mucus from the cervix and surrounding area using the large individually wrapped cleaning swab Discard the swab.



Open the package that contains the pink-capped Xpert Swab Transport tube and individually wrapped collection swab. Open the collection swab wrapper by peeling the top of the wrapper



Align the small groove against the edge (rim) of the tube and break it off. If needed, gently rotate the shaft to complete the breakage. Discard the top part of the swab shaft.



Hold the swab in your hand, placing your thumb and forefinger in the middle of the swab shaft.



Insert the collection swab into the endocervical canal. Rotate the swab for 30 seconds in the endocervical canal. Withdraw the swab carefully.



Unscrew the cap from the transport tube. Immediately place the collection swab into the transport tube.



8

Re-cap the transport tube and tighten the cap securely.

Label the transport tube with the sample ID and date of collection, as required.

Specimen Collection, Transport and Storage

Swab samples transferred to Xpert[®] CT/NG swab transport tube

Cepheid catalog # SWAB/A-50

Ē

or Cepheid catalog # SWAB/G-50 kits

	Swab Samples	Validated Collection Tool	Temperature (°C)	Storage Time
	Endocervical collection Swab	SWAB/A-50 SWAB/G-50 kits	+2 °C	60 days
	Vaginal collection Swab	SWAB/A-50 SWAB/G-50 kits	+ <u>2</u> + <u>30</u> °C	60 days
3	Pharyngeal collection Swab	SWAB/G-50 kits	+2 •C	60 days
9	Rectal collection Swab	SWAB/G-50 kits	+ <u>2</u> •C	60 days





Cartridge Preparation



Cartridge Preparation – Urine or Swab



1



Take one Xpert CT/NG cartridge and the provided transfer pipette



Label the side of the cartridge with the same ID as the collection tube



Open the cartridge lid



Gently mix by inverting the transport tube 3-4 times



Pipette at least 1 mL of the sample using the provided pipette*



Slowly empty the pipette into the sample chamber of the cartridge



Close the lid firmly. Start the test within the time frame specified in the package insert.



Adapted from document 301-1811



Run a Test



Scan cartridge barcode message



By default, do not click on Manual Entry or Cancel

Scan Cartridge Barcode

the beep

Patient ID patientid Sample ID sampleid Order Test - Assay

Cartridge barcode is successfully scanned when you hear

First Name



Scan the cartridge





Place the cartridge on the conveyor within 30 minutes of adding the sample.

"For complete details on how to run a test, refer to the Package Insert and the GeneXpert[®] Dx or Xpertise[™] Operator Manuals. 29 © Cepheid CE-IVD and US-IVD. For *in vitro* diagnostic use



Create a Test on GeneXpert[®] Dx Software



Ē

Create a Test on Xpertise[™] Software – Assay Selection





Create a Test on Xpertise[™] Software – Test Information

		Order Test - Test Information	
	Patient ID		
	patientid		
6 Review and complete the test information ——	Sample ID		
	sampleid		
	Last Name	First Nam	e
	patient	id	
	Assay*		Version*
	Xpert CT_NG		3
	Reagent Lot ID*	Cartridge S/N*	
	12102	282769448	
	Expiration Date*	Priority	
	2018/11/04	Normal	
	Test Type		
	Specimen	-	
	Sample Type	Other Sample Type	
	Other	▼	
	Notes		
7 Click on SUBMIT			
	-		SUBMIT 😡

8 Place the cartridge on the conveyor belt





Ē

Ē

Automated Xpert[®] Protocol



Quality Controls



Cepheid

Cepheid Control Strategy

Instrument System Control – Check Status

- System control checks the optics, temperature of the module and mechanical integrity of each cartridge.
- If the system controls fail, an ERROR test result will be reported.

Assay Quality Controls

- Each Xpert cartridge is a self-contained test device
- Cepheid designed specific molecular methods to include internal controls that enable the system to detect specific failure modes within each cartridge
 - Sample Adequacy Control (SAC)
 - Specimen Processing Control (SPC)
 - Probe Check Controls (PCC)



Internal Quality Controls

Sample Adequacy Control (SAC)

• Verifies that human cells are present in the sample

Probe Check Controls (PCC)

- Before the PCR step, fluorescence signal is measured on all probes and compared with pre-established factory settings to monitor
 - bead rehydration
- probe integrity
- reaction tube filling
- dye stability

Sample Processing Controls (SPC)

- Genomic DNA of Bacillus globigii in each cartridge
 - Verifies adequate sample processing
 - Verifies lysis, presence of the organism and detects PCR inhibition
 - Must be positive in a negative sample
 - Can be positive or negative in a positive sample



Commercially Available External Controls

Part Number	Description	Configuration	Storage
NATCT(434)-6MC	CT positive control	1 mL x 6 vials	2-8°C
NATNG-6MC	NG positive control	1 mL x 6 vials	2-8°C
NATCT/NGNEG-6MC	CT and NG Negative controls	1 mL x 6 vials	2-8°C
	http://www.zeptometrix.com		

- 1. Invert the control 3 to 4 times.
- 2. Open the cartridge lid.
- 3. Using a clean transfer pipette, fill the transfer pipette above the mark on the pipette shaft.
- 4. Ensure the pipette is filled with no air bubbles present.
- 5. Empty the contents of the pipette into the sample chamber with large opening in the cartridge.
- 6. Close cartridge lid.
- External controls should be used in accordance with local, state accrediting organizations, as applicable
- NATtrol[™] products are Research Use Only and not for in-vitro diagnostic use.



Result Interpretation

-

-

6

6

0

Cepheid

-

Result Interpretation Algorithm

F



Xpert[®] CT/NG – All possible results

Result displayed	CT1	NG2	NG4	SPC	SAC
CT DETECTED	_	_	_	±/	±/
NG DETECTED	т	т	т	+/-	+/-
CT DETECTED	1	1		+/	+/
NG NOT DETECTED	т	т	-	+/-	+/-
CT DETECTED	_			+/	+/
NG NOT DETECTED	т	-	Ŧ	±/-	+/-
CT NOT DETECTED		_		±/	±/
NG DETECTED	-	т	I	• 7 -	+/-
CT NOT DETECTED			+	+/-	±/
NG NOT DETECTED	-	-			+/-
CT NOT DETECTED					+
NG NOT DETECTED	-	-	-	Ŧ	т
INVALID	-	-	-	-	+/-
INVALID	_	-	-	+/-	-

Cepheid

F

CT DETECTED NG DETECTED

lyto .						
ne	Ct	EndPt		Analyte Result		Probe Check Result
CT1	30.2		261		POS	PASS
NG2	27.1		436		POS	PASS
NG4	26.3		531		POS	PASS
SAC	18.0		308		NA NA	PASS
nn+						Legend
						Image: Second state NG2; Primary Image: Second state NG4; Primary Image: Second state SAC; Primary Image: Second state SPC; Primary
00						
00+	/					
0 10	20 01	30 ycles		40		
	CT1 NG2 NG4 SAC SPC	CT1 30.2 NG2 27.1 NG4 26.3 SAC 18.0 SPC 33.6	CT1 30.2 NG2 27.1 NG4 26.3 SAC 18.0 SPC 33.6	CT1 30.2 261 NG2 27.1 436 NG4 26.3 531 SAC 18.0 308 SPC 33.6 219	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	CT1 30.2 261 NG2 27.1 436 POS SAC 18.0 308 SAC 18.0 308 NA SPC 33.6 219 00 00 00 00 00 00 00 00 00 0

- SAC: NA (not applicable)
 - SAC is ignored because a target amplification occurred
- SPC: NA (not applicable)
 - SPC is ignored because a target amplification occurred
- Probe Check: PASS





⁻ The targets CT1, NG2 and NG4 are detected and the Ct values are within the valid range

CT DETECTED; NG NOT DETECTED



- The target CT1 is detected and the Ct value is within the valid range
- None of the NG targets are detected
- SAC: NA (not applicable)
 - SAC is ignored because an amplification occurred
- SPC: NA (not applicable)
 - SPC is ignored because the CT1 target amplification occurred
- Probe Check: PASS





CT NOT DETECTED; NG DETECTED

CT NOT DETECTED; NG DETECTED



- The targets NG2 and NG4 are detected and the Ct values are within the valid range
- The target CT1 is not detected
- SAC: NA (not applicable)
 - SAC is ignored because the NG target amplification occurred
- SPC: NA (not applicable)
 - SPC is ignored because the NG target amplification occurred
- Probe Check: PASS



CT NOT DETECTED; NG NOT DETECTED



- The targets CT1, NG2 and NG4 are NOT detected
- SAC: PASS
 - SAC has a Ct value within the valid range
- SPC: PASS
 - SPC has a Ct value within the valid range
- Probe Check: PASS



CT NOT DETECTED NG NOT DETECTED

Troubleshooting



Factors That Negatively Affect Results

- Improper specimen collection
 - The number of organisms in the specimen is below the detection limit of the test
 - Performance with other collection devices and specimen types has not been assessed
- Improper transport or storage of collected specimen
 - Storage and transport conditions are specimen specific
 - Refer to the Package Insert for the appropriate handling instructions
- Improper testing procedure
 - Modification to the testing procedures may alter the performance of the test.
 - Careful compliance with the package insert is necessary to avoid erroneous results







- Presence or absence of the CT1 and NG2/4 targets can not be determined
- SAC: PASS
 - SPC has a Ct value within the valid range
- SPC: FAIL
 - SAC Ct value is not within the valid range
- Probe Check: PASS



F



- Presence or absence of the CT1 and NG2/4 targets can not be determined
- SAC: FAIL
 - SAC Ct value is not within the valid range
- SPC: PASS
 - SPC has a Ct value within the valid range
- Probe Check: PASS



INVALID

Ē



- Presence or absence of the CT1 and NG2/4 targets can not be determined
- SAC: FAIL
 - SPC Ct value is not within the valid range
- SPC: FAIL
 - SAC Ct value is not within the valid range
- Probe Check: PASS





- INVALID result with failing SPC and/or SAC <u>Origin(s)</u>
 - PCR was inhibited due to interfering substances
 - Inadequate sample was used
 - Improper specimen storage/collection/preparation
 - Improper kit storage conditions

Solution(s)

- Use the correct specimen type
- Check the sample quality (Blood, Mucin, topical medication...)
- Follow recommended instructions on sample collection, preparation and storage
- Check kit storage conditions and shelf life
- Collect a new sample when necessary and retest





- INVALID result with failing SAC only <u>Origin(s)</u>
 - Inadequate sample was used
 - Improper specimen collection
 - Improper sample storage or preparation
 - Improper kit storage conditions

Solution(s)

- Use the correct specimen type
- Check the collection: Urine first catch must be collected to ensure a proper epithelial cell concentration – Proper swabbing must be performed (according to illustrated collection instructions)
- Follow recommended instructions on sample collection, preparation and storage
- Check kit storage conditions and shelf life
- Collect a new sample in the appropriate conditions, when necessary, and retest





A A	Test Result	Analyte Result	Detail	Errors	History	Support
	Assay Name	Xpert CT_NG	Version	3		
	Test Result	ERROR				
	For In Vitro Dia	agnostic Use Only.				

- The Test result tab displays "ERROR"
- The error code and description can be found in the "Errors" Tab
- The test must be re-run, after corrective actions







Test Result NO RESULT
For In Vitro Diagnostic Use Only.

NO RESULT

- Test could not be completed and insufficient data was collected

ORIGIN(S)

- Power failure during test
- "Stop Test" function was used.
- Computer freeze or crash during test



Re-test Procedure

Discard used cartridge

Follow your institution's safety guidelines for disposal of cartridges

	0
P	
Provide Contraction	
-	-
The	100

9

Obtain the residual treated sample from either

- Swab Transport Reagent or
- Urine Transport Reagent tube

If the leftover sample volume is insufficient, or the retest continues to return an INVALID, ERROR, or NO RESULT, collect a new sample 3



Obtain a new cartridge

Label appropriately as retest on the new cartridge

Process the sample per the package insert





Run the test on the GeneXpert[®] System





-

-

Technical Assistance

- Before contacting Cepheid Technical Support, collect the following information:
 - Product name
 - Lot number
 - Serial number of the System
 - Error messages (if any)
 - Software version and, if applicable, Computer Service Tag number
- Log your complaint using the following link <u>http://www.cepheid.com/us/support</u>

Region	Telephone	Technical Support Email
US	+ 1 888 838 3222	techsupport@cepheid.com
Australia and New Zealand	+ 1800 130 821 + 0800 001 028	techsupportANZ@cepheid.com
Brazil and Latin America	+ 55 11 3524 8373	latamsupport@cepheid.com
China	+ 86 400 821 0728	techsupportchina@cepheid.com
France	+ 33 563 825 319	support@cepheideurope.com
Germany	+ 49 69 710 480 480	support@cepheideurope.com
India, Bangladesh, Bhutan, Nepal and Sri Lanka	+ 91 11 48353010	techsupportindia@cepheid.com
Italy	+ 39 800 902 567	support@cepheideurope.com
South Africa	+ 27 861 22 76 35	support@cepheideurope.com
United Kingdom	+ 44 3303 332 533	support@cepheideurope.com
Belgium, Luxembourg and Netherlands	+33 563 825 3319	support@cepheideurope.com
Other European, Middle East,	+ 33 563 825 319	aunnart@aanhaidaurana.aam
and African countries	+ 971 4 253 3218	support@ceprieldeurope.com
Other countries not listed	+1 408 400 8495	techsupport@cepheid.com



Thank You.

Xpert®

Cepheid.

GeneXpert

www.Cepheid.com

