



For use with GeneXpert® System with Touchscreen



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R_{only} IVD In Vitro Diagnostic Medical Device

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See Revision History for a description of changes.

1		
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	-	

Getting Started	5
Product Information	
Proprietary Name	
Common or Usual Name	
Intended Use, Summary, and Principle of Procedure	5
Intended Use	5
Summary and Explanation	5
Principle of the Procedure	_

	Reagents, Instruments, and Materials	7
\smile	Reagents	
	• Materials Provided	7
	Materials Required but Not Provided	
	Materials Available but Not Provided	
	Warnings and Precautions	8
	General	
	Specimen Collection	8
	Test/Reagent	8
	Chemical Hazards, Storage and Handling	10
	Chemical Hazards	10
	Storage and Handling	10

	Specimen Collection, Testing, and Results	
$\mathbf{}$	Specimen Collection	
	· Specimen Collection, Transport and Storage	
	Procedure	
	Preparing the Cartridge	13
	Starting the Test: GeneXpert System with Touchscreen	
	Viewing Results: GeneXpert System with Touchscreen	15
	Quality Control	15
	Built-in Quality Controls	
	External Controls	16
	Results	16
	Retests	17
	Reasons to Repeat the Test	
	Retest Procedure	
	Limitations	18
	Limitations of the Procedure	
	Expected Values	19

Specific Performance Characteristics	21
Clinical Performance	
Analytical Performance	23
Analytical Sensitivity (Limit of Detection)	23
Analytical Reactivity (Inclusivity)	24
Analytical Specificity (Cross-Reactivity and Competitive Interference)	24
Interfering Substances Study	
Carry-Over Contamination Study	
Reproducibility	
Instrument System Precision	

?

Appendix	
Bibliography	
Cepheid Headquarters Locations	
Technical Assistance	
Table of Symbols	41
Revision History	41



Product Information

Proprietary Name

Xpert[®] TV

Common or Usual Name

Xpert TV Test

Intended Use, Summary, and Principle of Procedure

Intended Use

The Xpert TV test, performed on the GeneXpert[®] Instrument Systems, is a qualitative *in vitro* diagnostic test for the detection of *Trichomonas vaginalis* genomic DNA. The test utilizes automated real-time polymerase chain reaction (PCR) to detect *Trichomonas vaginalis* genomic DNA. The Xpert TV test uses female and male urine specimens, endocervical swab specimens, or patient-collected vaginal swab specimens (collected in a clinical setting). The Xpert TV test is intended to aid in the diagnosis of trichomoniasis in symptomatic or asymptomatic individuals.

Summary and Explanation

The protozoan *Trichomonas vaginalis* is responsible for trichomoniasis, which is a common sexually transmitted infection that can infect both men and women. There are 7.4 million cases of trichomoniasis annually in the United States. Trichomoniasis infections can be symptomatic or asymptomatic.¹

In women, trichomoniasis is one of a range of conditions that comprise vaginal discharge. Symptoms in females can include itching, burning, redness, or soreness of the genitals, unusual odor, discomfort with urination, or a thin clear, white, yellow, or green discharge.² In men, trichomoniasis may cause non-gonococcal urethritis (NGU). Symptoms in males can include itching or burning inside the penis, burning after ejaculation or urination, or penile discharge.^{2,3}



Principle of the Procedure

The Xpert TV Test is an automated *in vitro* diagnostic test for qualitative detection of *Trichomonas vaginalis* (TV). The test is performed on Cepheid GeneXpert Instrument Systems.

The GeneXpert Instrument Systems automate and integrate sample preparation, nucleic acid extraction and amplification, and detection of the target sequence in simple or complex samples using reverse transcriptase polymerase chain reaction (RT-PCR) and/or real-time PCR tests. The systems consist of an instrument, personal computer, and preloaded software for running tests and viewing the results. The systems require the use of single-use disposable GeneXpert cartridges that hold the real-time PCR reagents and host the reverse transcriptase PCR and real-time PCR processes. Because the cartridges are self-contained, cross-contamination between samples is minimized. For a full description of the system, see the relevant system operator manual.

The Xpert TV Test includes reagents for the detection of *Trichomonas vaginalis*. The Xpert TV Test is designed for use with the following specimens collected from symptomatic and asymptomatic individuals: first-catch female and male urine, endocervical and vaginal swab specimens. The urine transport reagent and swab transport reagent are designed to preserve patient specimens during transport to the laboratory for analysis with Xpert TV Test and are included in the following specimen collection kits: Xpert Urine Specimen Collection Kit, the Xpert Swab Specimen Collection Kit, and the Xpert Vaginal/Endocervical Specimen Collection Kit.

A Sample Processing Control (SPC), a Sample Adequacy Control (SAC), and a Probe Check Control (PCC) are also included in the cartridge. The SPC is present to control for adequate processing of the target sample and to monitor the presence of inhibitors in the PCR reaction. The SAC reagents detect the presence of a single copy human gene and monitor whether the specimen contains human cells. The PCC verifies reagent rehydration, PCR tube filling in the cartridge, probe integrity, and dye stability.

An Early Assay Termination function provides positive results if target DNA reaches a predetermined threshold before the full 45 PCR cycles have been completed. When TV levels are high enough to generate very early Cts, neither the SAC nor SPC amplification curves will be seen and their results will not be reported.

Reagents, Instruments, and Materials

Reagents

Materials Provided

The Xpert TV Test kit (GXTV-10) contains sufficient reagents to process 10 specimens or quality control samples. The kit contains the following:

Xpert TV Test cartridges with integrated reaction tubes	10
• Bead 1, Bead 2, and Bead 3 (freeze-dried)	1 of each per cartridge
• Lysis Reagent (Guanidinium thiocyanate)	1.6 mL per cartridge
• Sodium Hydoxide	0.4 mL per cartridge
• Wash Reagent	0.5 mL per cartridge
• Elution Reagent	2.0 mL per cartridge
• Binding Reagent	1.5 mL per cartridge
Transfer Pipettes (500 μL)	10
CD	1

- Assay Definition File (ADF)
- Instructions to import ADF into GeneXpert software

• Instructions for use (Package Insert)

Note Safety Data Sheets (SDS) are available at www.cepheid.com or www.cepheidinternational.com under the **SUPPORT** tab.

Note The bovine serum albumin (BSA) in the beads within this product was produced and manufactured exclusively from bovine plasma sourced in the United States. No ruminant protein or other animal protein was fed to the animals; the animals passed ante- and post-mortem testing. During processing, there was no mixing of the material with other animal materials.



Materials Required but Not Provided

- Primary samples must be collected and treated with the appropriate kit:
 - URINE/A-50: Xpert Urine Specimen Collection Kit
 - $\circ\,$ SWAB/A-50: Xpert Vaginal/Endocervical Swab Specimen Collection Kit
 - SWAB/G-50 or SWAB/G-50-US: Xpert Swab Specimen Collection Kit
- GeneXpert system with touchscreen: GeneXpert instrument, touchscreen unit with built-in scanner, Cepheid OS software version 2.0 or higher, and operator manual.

Materials Available but Not Provided

- ZeptoMetrix NATtrol[™] TV External Run Control (catalog # NATTVNEG-6MC) as negative control.
- ZeptoMetrix NATtrol[™] TV External Run Control (catalog # NATTVPOS-6MC) as positive control.
- Printer (If a printer is required, contact Cepheid Technical Support to arrange for the purchase of a recommended printer.)

Warnings and Precautions

General

- For *in vitro* diagnostic use.
- 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. Guidelines for specimen handling are available from the U.S. Centers for Disease Control and Prevention⁴ and the Clinical and Laboratory Standards Institute.⁵
- Follow your institution's safety procedures for working with chemicals and handling biological samples.
- Good laboratory practices and changing gloves between handling patient specimens are recommended to avoid contamination of specimens.

Specimen Collection

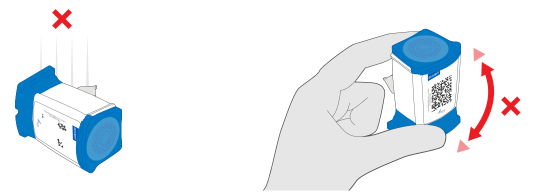
- For collection of endocervical swab specimens and patient-collected vaginal swab specimens, use only the Xpert Vaginal/Endocervical Specimen Collection Kit or Xpert Swab Specimen Collection Kit.
- For collection of urine specimens, use only the Xpert Urine Specimen Collection Kit with unpreserved (neat), first-catch urine.
- Under or over dispensing of urine into the Xpert Urine Transport Reagent tubes may affect test performance.
- Endocervical and patient-collected vaginal swab specimens must be collected and tested before the expiration date of the Xpert Swab Transport Reagent.
- Urine specimens must be collected and tested before the expiration date of the Xpert Urine Transport Reagent.

Test/Reagent

• Do not use a cartridge that has been dropped after removing from the kit or that has been shaken after the



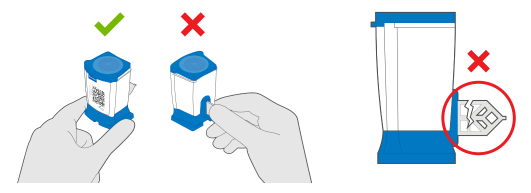
cartridge lid has been opened. Shaking or dropping the cartridge after opening the lid may yield false or non-determinate results.



• Do not place the sample ID label on the cartridge lid or on the barcode label.



• Hold the cartridge by the base. Do <u>not</u> touch the reaction tube at the rear of the cartridge as this could cause damage that would interfere with light passing through it during the test. Do not use a cartridge with a damaged reaction tube.



- Do not substitute Xpert TV Test reagents with other reagents.
- Do not open the Xpert TV Test cartridge lid until you are ready to add a sample during testing.
- Each single-use Xpert TV Test cartridge is used to process one test.
- Do not reuse processed cartridges.
- Each single-use disposable pipette is used to transfer one specimen.
- Do not use disposable pipettes more than one time.
- Do not test the endocervical or patient-collected vaginal specimens received in the laboratory without the swab present. A false negative test result may occur.
- Do not use a cartridge if it appears wet or if the lid seal appears to have been broken.



- CHANGE GLOVES if they come in contact with specimen or appear to be wet to avoid contaminating other specimens. Change gloves before leaving work area and upon entry into work area.
- Wear clean lab coats and gloves. Change gloves between processing each sample.
- In the event of a spill of specimens or controls, wear gloves and absorb the spill with paper towels. Then, thoroughly clean the contaminated area with a 1:10 dilution of freshly prepared household chlorine bleach. Final active chlorine concentration should be 0.5% regardless of the household bleach concentration in your country. Allow a minimum of two minutes of contact time. Ensure the work area is dry before using 70% denatured ethanol to remove bleach residue. Allow surface to dry completely before proceeding. Or, follow your institution's standard procedures for a contamination or spill event. For equipment, follow the manufacturer's recommendations for decontamination of equipment.
- Biological specimens, transfer devices and used cartridges should be considered capable of transmitting
 infectious agents requiring standard precautions. Follow your institution's environmental waste procedure
 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.⁶

Chemical Hazards, Storage and Handling

Chemical Hazards^{7,8}

UN GHS Hazard Pictogram: 🗘

Signal word: WARNING

- UN GHS Hazard Statements
- May be harmful if swallowed
- Causes mild skin irritation
- Causes serious eye irritation
- UN GHS Precautionary Statements
 - Prevention
 - Wash thoroughly after handling.
 - \circ Wear protective gloves/protection clothing/eye protection/face protection.
 - Response
 - Call a POISON CENTER or doctor/physician if you feel unwell.
 - $\,\circ\,$ If skin irritation occurs: Get medical advice/attention.
 - IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.
 - If eye irritation persists: Get medical advice/attention.

Storage and Handling

- Store the Xpert TV Test cartridges at 2–28°C.
- Do not open a cartridge until ready to perform testing.



- Use cartridges within 30 minutes after opening the cartridge lid.
- Do not use cartridges that have passed the expiration date.
- Do not use a cartridge that has leaked.
- Do not use any reagents that have become cloudy or discolored.

Specimen Collection, Testing, and Results

Specimen Collection

Specimen Collection, Transport and Storage

• Maintain proper storage conditions during specimen transport to ensure the integrity of the specimen. Specimen stability under shipping conditions other than those recommended has not been evaluated.

Refer to the appropriate specimen collection kit instructions for use for collection and transport instructions.

i Important Failure to store specimens as outlined in Table 1 through Table 3 may cause false negative results.

Specimen	Transport and Storage Temperature (°C)	Storage Time
Female and Male Urine	2–8 °C	4 days
	15–30 °C	4 hours

Table 1. Unprocessed Urine Specimen

Table 2. Urine Specimens in Xpert Urine Transport Reagent

Specimen	Transport and Storage Temperature (°C)	Storage Time
Female and Male Urine in Xpert Urine Transport Reagent	2–8 °C	28 days
remate and Mate Orine in Apert Orine Transport Reagent	15–30 °C	14 days

Table 3. Swab Specimens in Xpert Swab Transport Reagent

Specimen	Transport and Storage Temperature (°C)	Storage Time
Endocervical Swab in Xpert Swab Transport Reagent	2–30 °C	60 days
Vaginal Swab in Xpert Swab Transport Reagent	2–30 °C	60 days

Procedure



Preparing the Cartridge

(i) Important Start the test within 30 minutes of adding the sample to the cartridge.

To add the sample to the Xpert TV Test cartridge:

- 1. Obtain the following items:
 - Xpert TV Test cartridge
 - Transfer pipette (provided). Line on pipette indicates 500 µL fill volume.
 - Appropriately collected and labeled test sample in the Xpert Specimen Collection Kit transport reagent tube.
- 2. Inspect the test cartridge for damage. If damaged, do not use it.
- **3.** Open the cartridge lid.
- **4.** Gently invert the transport tube three to four times to ensure adequate mixing of sample and transport reagent.
- 5. Unwrap the transfer pipette.
- **6.** Remove the transport tube cap, compress the bulb of the transfer pipette, insert the pipette into the transport tube and release the bulb to fill the transfer pipette up to the mark (500 μ L) on the pipette shaft. See Figure 1. Ensure the pipette is filled with no air bubbles present.

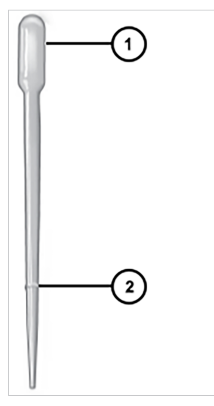


Figure 1 Transfer Pipette and Fill Mark

Number	Description
1	Bulb
2	Fill Transfer Pipette to the Mark



7. Empty the pipette's contents into the sample chamber of the cartridge. See Figure 2. Retain the remaining sample according to the conditions described in Table 2 and Table 3 in case a retest is required.



Figure 2 Xpert TV Test Cartridge (Top View)

8. Close the cartridge lid.

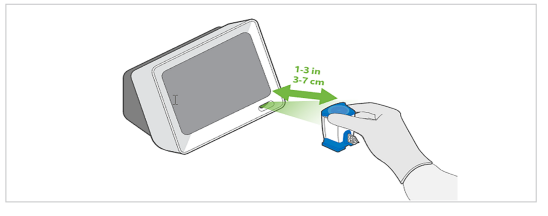
Starting the Test: GeneXpert System with Touchscreen

i) Important Before you start the test, make sure that:

- The system is running the correct Cepheid OS software version shown in section -Materials Required but Not Provided.
- The correct assay definition file is imported into the software.

Note The default workflow is shown. Your system administrator may alter the workflow.

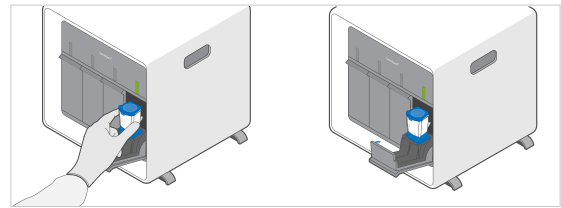
- **1.** Turn on GeneXpert system with touchscreen.
- 2. Log on to system software using your username and password.
- 3. On the Modules tab, touch Start Test.
- **4.** Follow onscreen prompts to create new test and enter patient and sample information.
- **5.** Scan or manually input the cartridge serial number. If scanning, hold the cartridge about 1-3 inches (3-7 cm) away from the scanner. The scanner projects a green crosshair, which you center on the barcode. Scanning is complete when you hear an audible beep. Touch **Continue**.



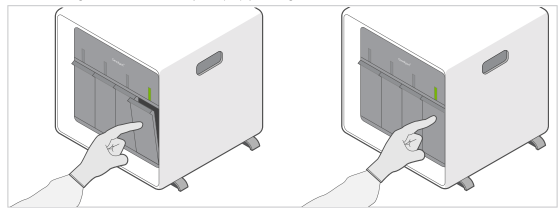
- 6. Select the desired test and touch **Continue**.
- 7. Watch the cartridge preparation video, if needed.



- **8.** On the Confirm screen, review all data and touch **Confirm**.
- **9.** Open the module door under flashing green light and insert the cartridge.



10. Close cartridge module door completely by pressing until it latches. The test starts.



- **11.** When the test completes, the **Results Summary** screen appears. Open the module door and remove cartridge.
- **12.** Dispose of used cartridge in appropriate waste container according to your institution's standard practices.

Viewing Results: GeneXpert System with Touchscreen

The GeneXpert system with touchscreen results screen will automatically interpret test results for you and clearly show them in the **View Results** window.

- 1. Tap Results.
- **2.** Tap the test to be viewed in the Results screen.
- 3. Click OK.
- **4.** To generate a PDF report file, touch **View Report**. More detailed instructions for viewing and uploading results are available in your system operator manual.

Quality Control



Built-in Quality Controls

Each test includes a Sample Processing Control (SPC), a Sample Adequacy Control (SAC), and Probe Check Control (PCC).

- Sample Processing Control (SPC): Ensures the sample was correctly processed. The SPC is included in each cartridge to verify adequate processing of the sample. The SPC verifies that binding and elution of *Trichomonas vaginalis* target DNA has occurred if the organism is present and verifies that the sample processing is adequate. Additionally, this control detects sample-associated inhibition of the real-time PCR test. The SPC should be positive in an analyte negative sample and can be negative or positive in an analyte positive sample. The SPC passes if it meets the validated acceptance criteria.
- Sample Adequacy Control (SAC): Verifies that the sample contains human cells or human DNA. This multiplex test includes primers and probes for the detection of a single copy human gene. The SAC signal is only to be considered in an analyte negative sample. A negative SAC indicates that no human cells are present in the sample due to insufficient mixing of the sample or because of an inadequately collected sample.
- **Probe Check Control (PCC):** Before the start of the PCR reaction, the GeneXpert System measures the fluorescence signal from the probes to monitor bead rehydration, reaction tube filling, probe integrity, and dye stability. The PCC passes if it meets the validated acceptance criteria.

External Controls

Positive and negative external controls should be used in accordance with local, state, and federal accrediting organizations' requirements as applicable.

Results

The results are interpreted automatically by the GeneXpert Instrument System from measured fluorescent signals and embedded calculation algorithms. Results are clearly shown on the Test Result tab of the View Results window. All possible Xpert TV Test results and their interpretation are shown in Table 4.

Result	Interpretation
TV DETECTED	 Trichomonas target DNA is detected. The Trichomonas target has a Ct within the valid range and a fluorescence endpoint above the threshold setting. SPC – Not applicable. SPC is ignored because the Trichomonas target amplification may compete with this control. SAC – Not applicable. SAC is ignored because the Trichomonas target amplification may compete with this control.' PCC – PASS. All probe check results pass.
TV NOT DETECTED	 Trichomonas target DNA is not detected. SPC meets acceptance criteria. Trichomonas target DNA is not detected. SPC – PASS. SPC has a Ct within the valid range and fluorescence endpoint above the threshold setting. SAC – PASS. SAC has a Ct within the valid range and a fluorescence endpoint above the threshold setting. PCC – PASS. All probe check results pass.



Result	Interpretation
	Presence or absence of <i>Trichomonas</i> target DNA cannot be determined. Repeat test according to the instructions in Retest Procedure.
	• SPC – FAIL. SPC Ct is not within valid range and the fluorescence endpoint is below the threshold setting.
	• SAC – PASS. SAC has a Ct within the valid range and fluorescence endpoint in the above threshold setting.
	• PCC – PASS. All probe check results pass.
	Or
INVALID	• SPC – PASS. SPC has a Ct within the valid range and fluorescence endpoint above the threshold setting.
	• SAC – FAIL. SAC Ct is not within valid range and fluorescence endpoint is below the threshold setting.
	• PCC – PASS. All probe check results pass.
	Or
	• SPC – FAIL. SPC Ct is not within valid range and fluorescence endpoint is below the threshold setting.
	• SAC – FAIL. SAC Ct is not within valid range and fluorescence endpoint is below the threshold setting.
	PCC – PASS. All probe check results pass.
	Presence or absence of <i>Trichomonas</i> target DNA cannot be determined. Repeat test according to the instructions in Retest Procedure.
	• TRICHOMONAS – NO RESULT
	• SPC – NO RESULT
ERROR	• SAC – NO RESULT
	• PCC – FAIL.* All or one of the probe check results fail.
	*If the probe check passed, the error is caused by the maximum pressure limit exceeding the acceptable range or by a system component failure.
	Presence or absence of <i>Trichomonas</i> target DNA cannot be determined. Repeat test according to the instructions in Retest Procedure. A NO RESULT indicates that insufficient data were collected. For example, the operator stopped a test that was in progress or a power failure occurred.
NO	TRICHOMONAS – NO RESULT
RESULT	• SPC – NO RESULT
	• SAC – NO RESULT
	• PCC – Not applicable

Retests

Reasons to Repeat the Test

If any of the following test results occur, repeat the test according to instructions in the Retest Procedure. Repeat the test using a new cartridge (do not re-use the cartridge).

- An **INVALID** result indicates that the SPC and/or the SAC failed. The sample was not properly processed, PCR was inhibited or the sample was not properly collected.
- An **ERROR** result indicates that the test failed possibly because the reaction tube was filled improperly, a reagent probe integrity problem was detected, pressure limits were exceeded, or a valve positioning error was detected.
- A **NO RESULT** indicates that insufficient data were collected. For example, the operator stopped a test that was in progress or a power failure occurred.



Retest Procedure

- Obtain the leftover sample from the Xpert Swab Transport Reagent Tube or Xpert Urine Transport Reagent Tube. Repeat the test with a new cartridge (do not re-use the cartridge). See the respective Starting the Test section.
- If the leftover sample volume is insufficient, or the retest continues to return an **INVALID**, **ERROR**, or **NO RESULT**, collect a new sample and repeat the test with a new cartridge.

Limitations

Limitations of the Procedure

- The Xpert TV Test has only been validated with the following specimen types, collected with the Xpert Vaginal/Endocervical Specimen Collection Kit, Xpert Swab Specimen Collection Kit, or the Xpert Urine Specimen Collection Kit:
 - $\circ \ {\rm Endocervical} \ {\rm swabs}$
 - Patient-collected vaginal swabs
 - \circ Female and male first-catch urine
- A negative test result does not exclude the possibility of infection because test results may be affected by improper specimen collection, technical error, sample mix-up, or because the number of organisms in the sample is below the limit of detection of the test.
- Careful compliance with the instructions in this instructions for use and in the Xpert Vaginal/Endocervical Specimen Collection Kit, Xpert Swab Specimen Collection Kit, and Xpert Urine Specimen Collection Kit instructions for use is necessary to avoid erroneous results.
- The Xpert TV Test has been validated using the procedures provided in this instructions for use only. Modifications to these procedures may alter the performance of the test.
- Because the detection of *Trichomonas vaginalis* is dependent on the organism's DNA present in the sample, reliable results are dependent on proper sample collection, handling, and storage.
- *Trichomonas tenax* was found to cross-react with the Xpert TV Test at levels above 1.0 x 10² cells/mL. *T. tenax* is a commensal of the oral cavity. See Xpert TV Analytical Specificity for details.
- With endocervical and patient-collected vaginal specimens, test interference may be observed in the presence of blood (> 60% v/v).
- As with many diagnostic tests, results from the Xpert TV Test should be interpreted in conjunction with other laboratory and clinical data available to the clinician.
- The patient-collected vaginal swab specimens are an option for screening women when a pelvic exam is not otherwise indicated.
- The Xpert TV Test has not been validated for use with vaginal swab specimens collected by patients at home. The patient collected vaginal swab specimen application is limited to healthcare facilities where support/counseling is available to explain procedures and precautions.
- The Xpert TV Test provides qualitative results. No correlation can be drawn between the magnitude of the Ct value and the number of cells in an infected sample.
- The Xpert TV Test should not be used for the evaluation of suspected sexual abuse or for other medico-legal indications.
- The predictive value of a test depends on the prevalence of the disease in any particular population. See Table 5 for hypothetical predictive values when testing varied populations.



- Mutations or nucleotide polymorphisms in primer or probe binding regions may affect detection of new or unknown *Trichomonas vaginalis* variants resulting in a false negative result.
- Xpert TV Test performance has not been evaluated in pregnant women, or in patients with a history of hysterectomy.
- Xpert TV Test performance has not been evaluated in patients less than 18 years of age or older than 78 years of age.

Expected Values

The prevalence of infection with *Trichomonas vaginalis* in patient populations depends on risk factors such as age, gender, the presence or absence of symptoms, the type of clinic, and the sensitivity of the test used to detect infections. During the clinical evaluation of the Xpert TV Test, the observed *Trichomonas vaginalis* prevalence rate in females was 10.3% and in males was 2.7%.

The estimated positive predictive value (PPV) and negative predictive value (NPV) of the Xpert TV Test across different hypothetical prevalence rates are shown for each specimen type in Table 5. These calculations are based on the overall estimated sensitivity and specificity observed for each specimen type during the Xpert TV multi-center clinical study (Table 6).

The overall sensitivity and specificity for male urine (UR-M) were 89.6% and 99.3%, respectively. The overall sensitivity and specificity for female urine (UR-F) were 98.4% and 99.7%, respectively. In patient-collected vaginal swab specimens (PC-VS), the overall sensitivity and specificity were 96.4% and 99.6%, respectively. For endocervical swabs (ES), the overall sensitivity and specificity were 98.9% and 98.9%, respectively.

Specimen Type	Prevalence (%)	PPV (%)	NPV (%)
	1	56.7%	99.9%
	2	72.6%	99.8%
	5	87.2%	99.5%
Male UR	10	93.5%	98.9%
Wate OK	12	94.6%	98.6%
	15	95.8%	98.2%
	20	97.0%	97.4%
	25	97.7%	96.6%
	1	76.2%	100.0%
	2	86.6%	100.0%
	5	94.3%	99.9%
Female UR	10	97.2%	99.8%
remale OK	12	97.7%	99.8%
	15	98.2%	99.7%
	20	98.8%	99.6%
	25	99.1%	99.5%
	1	69.0%	100.0%
PC-VS	2	81.8%	99.9%
	5	92.1%	99.8%

Table 5. Hypothetical PPV and NPV of the Xpert TV Test by Specimen Type



Specimen Type	Prevalence (%)	PPV (%)	NPV (%)
	10	96.1%	99.6%
	12	96.8%	99.5%
	15	97.5%	99.4%
	20	98.2%	99.1%
	25	98.7%	98.8%
	1	47.4%	100.0%
	2	64.5%	100.0%
	5	82.4%	99.9%
ES	10	90.8%	99.9%
ES	12	92.4%	99.8%
	15	94.0%	99.8%
	20	95.7%	99.7%
	25	96.7%	99.6%

I Specific Performance Characteristics

Clinical Performance

Performance characteristics of the Xpert TV Test were determined in a multi-site prospective investigational study by comparing the results from the Xpert TV Test to a patient infected status (PIS) algorithm comprised of culture and validated bi-directional sequencing (primary sequencing) for male urine, or an FDA-cleared NAAT test and culture for female specimen types.

Study participants included consenting asymptomatic and symptomatic, sexually active males and females seen at locations including, but not limited to: OB/GYN, sexually transmitted disease (STD), and family planning clinics. The average age among eligible female study participants was 33.5 years (range = 18 to 78 years). The average age among eligible male study participants was 36.2 years (range = 16 to 78 years).

The study specimens consisted of prospectively collected male urine, female urine, endocervical swabs, and patient-collected vaginal swabs (collected in a clinical setting). Clinician-collected vaginal swabs were collected for testing by the reference NAAT test and culture. Samples were collected from 17 clinical sites and tested at 11 sites. Reference testing was performed at 3 central laboratories.

A study participant was considered to be infected by PIS if either of the two reference test results were positive. The subject was considered to be not infected by PIS when both reference test results were negative.

Performance of the Xpert TV Test was calculated relative to the PIS for each of the three female specimen types (endocervical swabs, patient-collected vaginal swabs and urine) or to the PIS for male urine, respectively.

Specimens with discrepant results between the Xpert TV Test and the PIS were analyzed by validated bidirectional Sanger sequencing and results are footnoted in Table 6 for informational purposes only.

Among the 10,017 tests performed, 190 had initial **ERROR**, **INVALID**, or **NO RESULT** outcomes (1.90%, 95% CI 1.65-2.18). Of those, 167 specimens yielded valid results upon repeat test (7 specimens were not retested). The overall valid reporting rate of the test was 99.8% (9994/10,017).

Results of the Xpert TV Test were compared to the PIS for determination of sensitivity, specificity, and predictive values. Sensitivity and specificity for TV by specimen type and symptom status are presented in Table 6.



Sample Type	Status	Total (n)	Sens	95% CI	Spec	95% CI	Prev (%)	PPV (%)	NPV (%)
	Symp	685	100% (71/71)	94.9%-100%	98.5% (605/614)	97.2%-99.3%	10.4%	88.8%	100%
ES	Asymp	1114	98.1% (104/106)	93.4%-99.8%	99.1% (999/1008)	98.3%-99.6%	9.5%	92.0%	99.8%
	Overall	1799	98.9% (175/177) ^a	96.0%-99.9%	98.9% (1604/1622) ^b	98.3%-99.3%	9.8%	90.7%	99.9%
	Difference	P-Value	P=0.517	-0.70%, 4.48%	P=0.331	-1.69%, 0.54%			1
	Symp	682	98.6% (73/74)	92.7%-100%	99.5% (605/608)	98.6%-99.9%	10.9%	96.1%	99.8%
PC-VS	Asymp	1109	95.0% (113/119)	89.3%-98.1%	99.6% (986/990)	99.0%-99.9%	10.7%	96.6%	99.4%
	Overall	1791	96.4% (186/193) ^c	92.7%-98.5%	99.6% (1591/1598) ^d	99.1%-99.8%	10.8%	96.4%	99.6%
	Difference	P-Value	P=0.254	-1.04%, 8.42%	P=1.000	-0.77%, 0.59%			
	Symp	688	98.6% (71/72)	92.5%-100%	99.8% (615/616)	99.1%-100%	10.5%	98.6%	99.8%
UR-F	Asymp	1105	98.2% (109/111)	93.6%-99.8%	99.6% (990/994)	99.0.%-99.9%	10.0%	96.5%	99.8%
	Overall	1793	98.4% (180/183) ^e	95.3%-99.7%	99.7% (1605/1610) ^f	99.3%-99.9%	10.2%	97.3%	99.8%
	Difference	P-Value	P=1.000	-3.25%, 4.08%	P=0.655	-0.27%, 0.75%			1
UR-M	Symp	1088	87.5% (28/32)	71.9%-95.0%	99.8% (1054/1056)	99.3%-99.9%	2.9%	93.3%	99.6%
	Asymp	3523	90.3% (84/93)	82.6%-94.8%	99.2% (3401/3430)	98.8%-99.4%	2.6%	74.3%	99.7%
	Overall	4611	89.6% (112/125) ^g	83.0%-93.8%	99.3% (4455/4486) ^h	99.0%-99.5%	2.7%	78.3%	99.7%
	Difference	P-Value	P=0.738	-15.8%, 10.1%	P=0.020	0.25%, 1.06%			

Table 6. Xpert TV vs PIS by Symptomatic Status

a. Testing results by sequencing: 1 of 2 FN was TV positive; 1 of 2 was TV negative.

b. Testing results by sequencing: 8 of 18 FP were TV positive; 10 of 18 were TV negative.

- c. Testing results by sequencing: 3 of 7 FN were TV positive; 4 of 7 were TV negative.
- d. Testing results by sequencing: 5 of 7 FP were TV positive; 2 of 7 were TV negative.
- e. Testing results by sequencing: 3 of 3 FN were TV negative.
- f. Testing results by sequencing: 5 of 5 FP were TV negative.
- g. Testing results by secondary sequencing: 9 of 13 false negatives were TV negative; 4 of 13 were TV positive.
- h. Testing results by secondary sequencing: 27 of 31 false positives were TV positive; 4 of 31 were TV negative.

Cycle Threshold (Ct) Frequency Distribution

Patient-collected vaginal swabs, endocervical swabs and urine specimens were collected from 1867 females and urine specimens were collected from 4626 males at 17 collection sites in the US. The frequency distribution of Xpert TV Test positive results for the 197 *Trichomonas vaginalis* infected female study subjects and 125 *Trichomonas vaginalis* infected male study subjects are shown in Figure 3.

swab, UR-F=female urine, UR-M=male urine



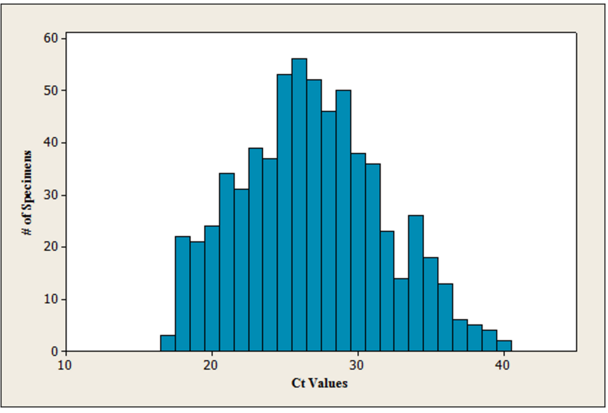


Figure 3 Ct Distribution of Patients Designated as Positive for TV Based on PIS Algorithm

Analytical Performance

Analytical Sensitivity (Limit of Detection)

The analytical sensitivity or limit of detection (LoD) of the Xpert TV Test was assessed using two *Trichomonas* vaginalis strains, one metronidazole susceptible (*T. vaginalis* $ATCC^{$ [®] 30001^M), and one metronidazole

resistant (*T. vaginalis* ATCC[®] 30238[™]). The strains were tested individually in clinical *T. vaginalis*-negative pooled urine matrix in Cepheid Xpert Urine Transport Reagent and clinical *T. vaginalis*-negative pooled vaginal swab matrix (VS) in Cepheid Xpert Swab Transport Reagent.

T. vaginalis was cultured and incubated at 35° C. Visual examination of the cultures for white precipitate (indicating growth) was conducted every 24 hours for 3 to 5 days. Cell pellets were resuspended in growth medium and enumerated visually using light microscopy. The concentration of isolates was expressed as the number of cells per milliliter (cells/mL). Cultures were diluted in culture medium to 1×10^4 cells/mL and stored at -20 °C. Cells were thawed on ice for use in the study.

The LoD was estimated by testing replicates of 20 at five concentrations for each strain and sample type over three days. The LoD for each strain was estimated by probit analysis. The claimed LoDs were confirmed by analyzing at least 20 replicates with *T. vaginalis* cells diluted to the estimated LoD concentrations. The LoD is defined as the lowest number of cells/mL that can be reproducibly distinguished from negative samples with 95% confidence or the lowest concentration at which 19 of 20 replicates were positive. The study was performed with two different lots of Xpert TV reagents and the claimed LoD for each strain is the higher of the two determinations (Table 7). The claimed LoD for *T. vaginalis* strains ATCC 30001 and ATCC 30238 in vaginal swab matrix is 2 cells/mL. The claimed LoD for *T. vaginalis* strain ATCC 30001 in urine matrix is 3 cells/mL. The claimed LoD for *T. vaginalis* in urine matrix is 2 cells/mL.



Trichomonas vaginalis	LoD Estimates by Probit Analysis (cells/mL)		Verified LoD	Verification	Mean	Mean	Mean	LoD Claim
strain and matrix	Reagent Lot 1	Reagent Lot 2	(cells/mL)	(Positives/20)	TV Ct	SAC Ct	SPC Ct	(cells/mL)
ATCC 30001 in Vaginal Swab	2.0	1.6	2.0	20/20	39.1	21.4	33.9	2
ATCC 30238 in Vaginal Swab	1.7	2.1	2.1	20/20	37.5	21.4	33.7	2
ATCC 30001 in Urine	2.2	2.5	2.5	20/20	38.2	29.3	34.1	3
ATCC 30238 in Urine	2.1	1.7	2.1	20/20	38.2	29.2	33.8	2

Table 7. LoD of Two T. vaginalis Strains in Pooled Vaginal Swab Matrix and Urine Matrix
Table 1. Lob of 1. Vaginalis Strains in 1. oolea Vaginal Swab Matrix and Ornie Matrix

Analytical Reactivity (Inclusivity)

The analytical inclusivity of the Xpert TV Test was evaluated by testing 17 *T. vaginalis* strains diluted in either negative pooled vaginal swab matrix in Cepheid Xpert Swab Transport Reagent or negative pooled urine in Cepheid Xpert Urine Transport Reagent. All *T. vaginalis* strains were tested in triplicate at a concentration of 3X the analytical LoD for the respective specimen type (6 cells/mL for vaginal swabs and 7.5 cells/mL for urine). All strains tested were reported as **TV DETECTED**. Results are shown in Table 8. Positive and negative controls were included in the study. The inclusivity for the 17 *T. vaginalis* strains tested was 100%.

Isolate ATCC #	Isolation Source	Results Vaginal Swab	Results Urine
30001	Vaginal exudate	TV DETECTED	TV DETECTED
30184	Vaginal swab	TV DETECTED	TV DETECTED
30187	Endocervical swab	TV DETECTED	TV DETECTED
30188	Vagina	TV DETECTED	TV DETECTED
30236	Endocervical swab	TV DETECTED	TV DETECTED
30240	Vaginal pool	TV DETECTED	TV DETECTED
30245	Vaginal and Endocervical material	TV DETECTED	TV DETECTED
30247	Vagina	TV DETECTED	TV DETECTED
50138	human	TV DETECTED	TV DETECTED
50139	human	TV DETECTED	TV DETECTED
50141	human	TV DETECTED	TV DETECTED
50143	human	TV DETECTED	TV DETECTED
50147	human	TV DETECTED	TV DETECTED
50167	Vagina	TV DETECTED	TV DETECTED
50183	Prostatic fluid	TV DETECTED	TV DETECTED
PRA-95	Vaginal exudate	TV DETECTED	TV DETECTED
PRA-98	human	TV DETECTED	TV DETECTED

Table 8. Analytical Reactivity (Inclusivity) of Xpert TV Test

Analytical Specificity (Cross-Reactivity and Competitive Interference)

A panel of 124 microorganisms, including bacteria, fungi, and viruses commonly found in the urogenital



tract, as well as other protozoans closely related to *T. vaginalis* were tested with the Xpert TV Test. The microorganisms were tested in the presence (competitive interference) and absence (cross-reactivity) of 3X LoD *T. vaginalis* ATCC 30001 cells. The microorganisms were seeded into either pooled *Trichomonas vaginalis*-negative urine matrix (patient urine added to Cepheid Urine Transport Reagent) or pooled *Trichomonas vaginalis*-negative vaginal swab matrix (vaginal swabs collected into Cepheid Swab Transport Reagent).

Each bacterial or fungal strain was tested at 1×10^6 CFU/mL or greater or at 1×10^6 genomes/mL. Viral strains were tested at 1×10^5 TCID₅₀/mL or 10^5 genomes/mL or greater. Protozoans were cultured in growth media, visually enumerated by light microscopy and tested at 1×10^5 cells/mL or greater or 10^5 genomes/mL. All microorganisms were tested in triplicate. Positive and negative controls were included in the study. One organism, *Trichomonas tenax*, demonstrated cross-reactivity (result of **TV DETECTED** in the absence of TV) at 1×10^5 cells/mL for the urine and vaginal swab matrix samples. *Trichomonas tenax* was subjected to repeat analysis at various other concentrations until a result of **TV NOT DETECTED** was obtained (at 1×10^2 cells/ mL). This is addressed in Limitations of the Procedure. For the other 123 microorganisms, all TV positive samples remained positive and all TV negative samples remained negative, indicating that there was no interference or cross-reactivity with the results of the Xpert TV Test for these microorganisms. Results are shown in Table 9 and Table 10 for urine and vaginal swab matrix, respectively.

	Xpert TV Test Result			
Microorganism	Concentration Tested ^a	Cross Reactivity (- T. vaginalis)	Competitive Interference (+ T. vaginalis)	
Achromobacter xerosis	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED	
Acinetobacter calcoaceticus	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED	
Acinetobacter lwoffii	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED	
Actinomyces israelii ^b	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED	
Actinomyces pyogenes	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED	
Aerococcus viridans	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED	
Aeromonas hydrophila	1 x 10 ⁷	TV NOT DETECTED	TV DETECTED	
Alcaligenes faecalis	1 x 10 7	TV NOT DETECTED	TV DETECTED	
Atopobium vaginae ^b	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED	
Bacillus subtilis	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED	
Bacteroides fragilis ^b	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED	
Bacteroides ureolyticus ^b	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED	
Bifidobacterium adolescentis ^b	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED	
Bifidobacterium brevi (breve) ^b	9 x 10 ⁶	TV NOT DETECTED	TV DETECTED	
Blastocystis hominis ^c	1 x 10 ^{5 d}	TV NOT DETECTED	TV DETECTED	
Branhamella catarrhalis	1 x 10 ⁷	TV NOT DETECTED	TV DETECTED	
Brevibacterium linens	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED	
Campylobacter jejuni	1 x 10 ⁷	TV NOT DETECTED	TV DETECTED	

Table 9. Analytical Specificity/Competitive Interference Determination for Xpert TV Test in Urine Matrix



	Xpert TV Test Result				
Microorganism	Concentration Tested ^a	Cross Reactivity	Competitive Interference		
	Concentration rested	(- T. vaginalis)	(+ T. vaginalis)		
Candida albicans ^e	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Candida glabrata ^e	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Candida parapsilosis ^e	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Candida tropicalis ^e	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Chlamydia trachomatis	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Chromobacterium violaceum	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Citrobacter freundii	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Clostridiodes difficile ^b	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Clostridium perfringens ^b	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Corynebacterium genitalium	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Corynebacterium xerosis	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Cryptococcus neoformans ^e	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Cryptosporidium parvum ^c	1 x 10 ^{5 d}	TV NOT DETECTED	TV DETECTED		
Cytomegalovirus ^f	5 x 10 ⁵	TV NOT DETECTED	TV DETECTED		
Deinococcus radiodurans	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Derxia gummosa	1 x 10 ^{6 d}	TV NOT DETECTED	TV DETECTED		
Eikenella corrodens	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Entamoeba histolytica ^c	1 x 10 ^{5 d}	TV NOT DETECTED	TV DETECTED		
Enterobacter aerogenes	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Enterobacter cloacae	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Enterococcus avium	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Enterococcus faecalis	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Enterococcus faecium	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Erysipelothrix rhusiopathiae	1 x 10 ⁷	TV NOT DETECTED	TV DETECTED		
Escherichia coli	1 x 10 ⁷	TV NOT DETECTED	TV DETECTED		
Flavobacterium meningosepticum	1 x 10 ⁷	TV NOT DETECTED	TV DETECTED		
Fusobacterium nucleatum ^b	4 × 10 ⁶	TV NOT DETECTED	TV DETECTED		
Gardnerella vaginalis	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Gemella haemolysans	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Giardia intestinalis ^c	1 x 10 ^{5 d}	TV NOT DETECTED	TV DETECTED		
Haemophilus ducreyi	1 x 10 ^{6 d}	TV NOT DETECTED	TV DETECTED		
Haemophilus influenzae	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Herpes simplex virus I ^f	1 x 10 ⁵	TV NOT DETECTED	TV DETECTED		



	Xpert TV Test Result					
Microorganism		Cross Reactivity	Competitive Interference			
	Concentration Tested ^a	(- T. vaginalis)	(+ T. vaginalis)			
Herpes simplex virus II ^f	1 x 10 ⁵	TV NOT DETECTED	TV DETECTED			
HIV-1 ^f	2 x 10 ⁵	TV NOT DETECTED	TV DETECTED			
Human papilloma virus 16 ^f	6 x 10 ⁵	TV NOT DETECTED	TV DETECTED			
Kingella dentrificans	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Kingella kingae	1 x 10 ⁷	TV NOT DETECTED	TV DETECTED			
Klebsiella oxytoca	1 x 10 ⁷	TV NOT DETECTED	TV DETECTED			
Klebsiella pneumoniae	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Lactobacillus acidophilus	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Lactobacillus brevis	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Lactobacillus crispatus	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Lactobacillus jensonii	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Lactobacillus lactis	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Lactobacillus vaginalis	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Legionella pneumophila	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Leuconostoc paramesenteroides	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Listeria monocytogenes	1 x 10 ⁷	TV NOT DETECTED	TV DETECTED			
Micrococcus luteus	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Mobiluncus curtisii ^b	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Moraxella lacunata	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Moraxella osloensis	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Morganella morganii	1 x 10 ⁷	TV NOT DETECTED	TV DETECTED			
Mycobacterium smegmatis	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Mycoplasma genitalium	1 x 10 ^{6 d}	TV NOT DETECTED	TV DETECTED			
Mycoplasma hominis	1 x 10 ^{6 d}	TV NOT DETECTED	TV DETECTED			
Neisseria cinerea	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Neisseria dentrificans	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Neisseria elongata	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Neisseria flava	9 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Neisseria flavescens	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Neisseria gonorrhoeae	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Neisseria lactamica	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Neisseria mucosa	9 x 10 ⁶	TV NOT DETECTED	TV DETECTED			
Neisseria perflava	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED			



	Xpert TV Test Result				
Microorganism		Cross Reactivity	Competitive Interference		
	Concentration Tested ^a	(- T. vaginalis)	(+ T. vaginalis)		
Neisseria polysaccharea	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Neisseria sicca	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Neisseria subflava	9 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Pantoea agglomerans	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Paracoccus denitrificans	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Pentatrichomonis hominis ^c	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Peptostreptococcus anaerobius ^b	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Peptostreptococcus productus ^b	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Plesiomonas shigelloides	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Prevotella bivia ^b	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Propionibacterium acnes ^b	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Proteus mirabilis	1 x 10 ⁷	TV NOT DETECTED	TV DETECTED		
Proteus vulgaris	1 x 10 ⁷	TV NOT DETECTED	TV DETECTED		
Providencia stuartii	1 x 10 ⁷	TV NOT DETECTED	TV DETECTED		
Pseudomonas aeruginosa	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Pseudomonas fluorescens	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Pseudomonas putida	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Rahnella aquatilis	1 x 10 ⁷	TV NOT DETECTED	TV DETECTED		
Rhodospirillum rubrum	1 x 10 ^{6 d}	TV NOT DETECTED	TV DETECTED		
Saccharomyces cerevisiae ^e	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Salmonella minnesota	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Salmonella typhimurium	1 x 10 ⁷	TV NOT DETECTED	TV DETECTED		
Serratia marcescens	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Staphylococcus aureus	9 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Staphylococcus epidermidis	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Staphylococcus saprophyticus	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Streptococcus agalactiae	1 x 10 7	TV NOT DETECTED	TV DETECTED		
Streptococcus bovis	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Streptococcus mitis	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Streptococcus mutans	2 x 10 ⁷	TV NOT DETECTED	TV DETECTED		
Streptococcus pneumoniae	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Streptococcus pyogenes	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED		
Streptococcus salivarius	9 x 10 ⁶	TV NOT DETECTED	TV DETECTED		



		Xpert TV Test Result								
Microorganism	Concentration Tested ^a	Cross Reactivity (- T. vaginalis)	Competitive Interference (+ T. vaginalis)							
Streptococcus sanguis	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED							
Streptomyces griseinus	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED							
Trichomonas tenax ^c	1 x 10 ⁵	TV DETECTED	TV DETECTED							
Trichomonas tenax ^c	1 x 10 ³	TV DETECTED	TV DETECTED							
Trichomonas tenax ^c	1 x 10 ²	TV NOT DETECTED	TV DETECTED							
Ureaplasma parvum	1 x 10 ^{6 d}	TV NOT DETECTED	TV DETECTED							
Ureaplasma urealyticum	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED							
Vibrio parahaemolyticus	9 x 10 ⁶	TV NOT DETECTED	TV DETECTED							
Yersinia enterocolitica	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED							

a. Tests run $\geq 10^{6}$ CFU/mL for bacteria and fungi, $\geq 10^{6}$ genomes/mL for yeast, $\geq 10^{5}$ TCID₅₀/mL or $\geq 10^{5}$ genomes/mL for viruses and $\geq 10^{5}$ cells/mL for protozoans.

- b. Anaerobic organism
- c. Protozoan
- d. Genome equivalents tested (DNA)
- e. Fungal organism
- f. Virus

Table 10. Analytical Specificity/Competitive Interference Determination for Xpert TV Test in Vaginal Swab Matrix

		Xpert TV Test Result					
Microorganism	Concentration Tested ^a	Cross-Reactivity (- T. vaginalis)	Competitive Interference (+ T. vaginalis)				
Achromobacter xerosis	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Acinetobacter calcoaceticus	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Acinetobacter lwoffii	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Actinomyces israelii ^b	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Actinomyces pyogenes	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Aerococcus viridans	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Aeromonas hydrophila	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Alcaligenes faecalis	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Atopobium vaginae ^b	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Bacillus subtilis	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Bacteroides fragilis ^b	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Bacteroides ureolyticus ^b	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED				



		Xpert TV Test Result					
Microorganism	Concentration Tested ^a	Cross-Reactivity	Competitive Interference				
		(- T. vaginalis)	(+ T. vaginalis)				
Bifidobacterium adolescentis ^b	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Bifidobacterium brevi (breve) ^b	9 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Blastocystis hominis ^c	1 x 10 ^{5 d}	TV NOT DETECTED	TV DETECTED				
Branhamella catarrhalis	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Brevibacterium linens	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Campylobacter jejuni	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Candida albicans ^e	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Candida glabrata ^e	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Candida parapsilosis ^e	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Candida tropicalis ^e	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Chlamydia trachomatis	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Chromobacterium violaceum	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Citrobacter freundii	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Clostridiodes difficile ^b	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Clostridium perfringens ^b	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Corynebacterium genitalium	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Corynebacterium xerosis	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Cryptococcus neoformans ^e	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Cryptosporidium parvum ^c	1 x 10 ^{5 d}	TV NOT DETECTED	TV DETECTED				
Cytomegalovirus ^f	5 x 10 ⁵	TV NOT DETECTED	TV DETECTED				
Deinococcus radiodurans	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Derxia gummosa	1 x 10 ^{6 d}	TV NOT DETECTED	TV DETECTED				
Eikenella corrodens	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Entamoeba histolytica ^c	1 x 10 ^{5 d}	TV NOT DETECTED	TV DETECTED				
Enterobacter aerogenes	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Enterobacter cloacae	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Enterococcus avium	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Enterococcus faecalis	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Enterococcus faecium	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Erysipelothrix rhusiopathiae	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Escherichia coli	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Flavobacterium meningosepticum	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Fusobacterium nucleatum ^b	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED				



		Xpert TV Test Result					
Microorganism	Concentration Tested ^a	Cross-Reactivity	Competitive Interference				
			(+ T. vaginalis)				
Gardnerella vaginalis	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Gemella haemolysans	9 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Giardia intestinalis ^c	1 x 10 ^{5 d}	TV NOT DETECTED	TV DETECTED				
Haemophilus ducreyi	1 x 10 ^{6 d}	TV NOT DETECTED	TV DETECTED				
Haemophilus influenzae	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Herpes simplex virus I ^f	1 x 10 ⁵	TV NOT DETECTED	TV DETECTED				
Herpes simplex virus II ^f	1 x 10 ⁵	TV NOT DETECTED	TV DETECTED				
HIV-1 ^f	2 x 10 ⁵	TV NOT DETECTED	TV DETECTED				
Human papilloma virus 16 ^f	6 x 10 ⁵	TV NOT DETECTED	TV DETECTED				
Kingella dentrificans	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Kingella kingae	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Klebsiella oxytoca	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Klebsiella pneumoniae	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Lactobacillus acidophilus	9 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Lactobacillus brevis	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Lactobacillus crispatus	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Lactobacillus jensonii	9 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Lactobacillus lactis	1 x 10 ⁷	TV NOT DETECTED	TV DETECTED				
Lactobacillus vaginalis	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Legionella pneumophila	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Leuconostoc paramesenteroides	9 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Listeria monocytogenes	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Micrococcus luteus	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Mobiluncus curtisii ^b	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Moraxella lacunata	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Moraxella osloensis	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Morganella morganii	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Mycobacterium smegmatis	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Mycoplasma genitalium	1 x 10 ^{6 d}	TV NOT DETECTED	TV DETECTED				
Mycoplasma hominis	1 × 10 ^{6 d}	TV NOT DETECTED	TV DETECTED				
Neisseria cinerea	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Neisseria dentrificans	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Neisseria elongata	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED				



		Xpert TV Test Result					
Microorganism	Concentration Tested ^a	Cross-Reactivity	Competitive Interference				
		(- T. vaginalis)	(+ T. vaginalis)				
Neisseria flava	9 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Neisseria flavescens	8 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Neisseria gonorrhoeae	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Neisseria lactamica	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Neisseria mucosa	9 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Neisseria perflava	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Neisseria polysaccharea	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Neisseria sicca	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Neisseria subflava	9 x 10 6	TV NOT DETECTED	TV DETECTED				
Pantoea agglomerans	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Paracoccus denitrificans	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Pentatrichomonis hominis ^c	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Peptostreptococcus anaerobius ^b	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Peptostreptococcus productus ^b	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Plesiomonas shigelloides	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Prevotella bivia ^b	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Propionibacterium acnes ^b	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Proteus mirabilis	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Proteus vulgaris	7 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Providencia stuartii	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Pseudomonas aeruginosa	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Pseudomonas fluorescens	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Pseudomonas putida	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Rahnella aquatilis	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Rhodospirillum rubrum	1 x 10 ^{6 d}	TV NOT DETECTED	TV DETECTED				
Saccharomyces cerevisiae ^e	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Salmonella minnesota	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Salmonella typhimurium	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Serratia marcescens	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Staphylococcus aureus	9 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Staphylococcus epidermidis	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Staphylococcus saprophyticus	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED				
Streptococcus agalactiae	6 x 10 ⁶	TV NOT DETECTED	TV DETECTED				



		Xpert TV Test Result						
Microorganism	Concentration Tested ^a	Cross-Reactivity (- T. vaginalis)	Competitive Interference (+ T. vaginalis)					
Streptococcus bovis	4 × 10 ⁶	TV NOT DETECTED	TV DETECTED					
Streptococcus mitis	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED					
Streptococcus mutans	5 x 10 ⁶	TV NOT DETECTED	TV DETECTED					
Streptococcus pneumoniae	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED					
Streptococcus pyogenes	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED					
Streptococcus salivarius	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED					
Streptococcus sanguis	2 x 10 ⁶	TV NOT DETECTED	TV DETECTED					
Streptomyces griseinus	4 x 10 ⁶	TV NOT DETECTED	TV DETECTED					
Trichomonas tenax ^c	1 x 10 ⁵	TV DETECTED	TV DETECTED					
Trichomonas tenax ^c	1 x 10 ³	TV DETECTED	TV DETECTED					
Trichomonas tenax ^c	1 x 10 ²	TV NOT DETECTED	TV DETECTED					
Ureaplasma parvum	1 x 10 ^{6 d}	TV NOT DETECTED	TV DETECTED					
Ureaplasma urealyticum	1 x 10 ⁶	TV NOT DETECTED	TV DETECTED					
Vibrio parahaemolyticus	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED					
Yersinia enterocolitica	3 x 10 ⁶	TV NOT DETECTED	TV DETECTED					

a. Tests run ≥ 106 CFU/mL for bacteria and fungi, $\geq 10^{6}$ genomes/mL for yeast, $\geq 10^{5}$ TCID50/mL or $\geq 10^{5}$ genomes/mL for viruses and $\geq 10^{5}$ cells/mL for protozoans.

b. Anaerobic organism

- c. Protozoan
- d. Fungal organism
- e. Virus

Additional three microorganisms, *Dientamoeba fragilis*, *Agrobacterium radiobacter*, and *Erwinia herbicola*, were not available for direct testing. An *in silico* analysis was conducted using the Basic Local Alignment Search Tool (BLAST) to compare the Xpert TV Test primer and probe sequences with all available sequences associated with these three microorganisms in the GenBank database. Available sequence data for *D. fragilis* was examined and showed a maximum of 7% homology to the Xpert TV primer and probe sequences. Available sequence data for *A. radiobacter* was examined and showed a maximum of 38% homology to the Xpert TV primer and probe sequences. Available sequence data for *A. radiobacter* was examined and showed a maximum of 38% homology to the Xpert TV primer and probe sequences. Available sequence data for *B. herbicola* was examined and showed a maximum of 10% homology to the Xpert TV primer and probe sequences.

Table 11. In silico Analytical Specificity Determination for Xpert TV Test

Strain	Accession Number	% Homology
Dientamoeba fragilis	KC967121.1	7%
Agrobacterium radiobacter	CP000629.1	38%
Erwinia herbicola	NG_035384.1	10%

Interfering Substances Study

The performance of the Xpert TV Test was evaluated with potentially interfering endogenous and exogenous substances that may be present in the urogenital tract.

All substances were tested in the presence and absence of 3X LoD *T. vaginalis* (ATCC strain 30001) to determine if there was interference with the Xpert TV Test. Substances were individually diluted into either pooled *Trichomonas vaginalis*-negative urine matrix (patient urine added to Cepheid Urine Transport Reagent) or pooled *Trichomonas vaginalis*-negative vaginal swab matrix (vaginal swabs collected into Cepheid Swab Transport Reagent). Positive and negative controls were included in the study.

For each interfering substance, eight replicates were tested for each set of samples (either *T. vaginalis* negative or *T. vaginalis* positive in clinical matrix). Table 12 and Table 13 show the substances that were tested, the test concentrations, and the matrix in which they were diluted. One substance, blood at > 60% v/v demonstrated interference (result of **TV NOT DETECTED** in the presence of TV) in the vaginal swab matrix samples. Blood was subjected to repeat analysis at various lower concentrations until a result of **TV DETECTED** was obtained (50% v/v). For the other conditions and substances tested, all TV positive samples remained positive and all TV negative samples remained negative, indicating that there was no interference causing false negative or false positive results with the Xpert TV Test for these substances.

Class/Substance	Active Ingredient	Concentration Tested				
Blood	Blood	0.3% v/v, 1% v/v				
Seminal Fluid	Seminal Fluid	5.0% v/v				
Mucus	Mucin	0.8% w/v				
	Acetylsalicylic Acid 500 mg	40 mg/mL				
Analassias 8 Antihistias	Acetaminophen	3.2 mg/mL				
Analgesics & Antibiotics	Azithromycin	1.8 mg/mL				
	Doxycycline	3.6 mg/mL				
OTC Deodorant & Powders	PEG-20; PEG-32; PEG-20 Stearate	0.25% w/v				
OTC Deodorant & Powders	Nanoxynol-9	0.25% w/v				
Albumin	BSA	10 mg/mL				
Glucose	Glucose	10 mg/mL				
Bilirubin	Bilirubin	1 mg/mL				
Acidic Urine (pH 4.0)	Urine + N-Acetyl-L-Cysteine	pH 4.0				
Alkaline Urine (pH 9.0)	Urine + Ammonium Citrate	pH 9.0				
Leukocytes	Leukocytes	10 ⁵ cells/mL				
Intravaginal Hormones	Progesterone; Estradiol	7 mg/mL Progesterone + 0.07 mg/mL Beta Estradiol				

Table 12. Potentially Interfering Substances in Urine Samples

Table 13. Potentially Interfering Substances in Swab Samples

Class/Substance	Active Ingredient	Concentration Tested			
Blood ^a	Blood	10%, 50%, 60% v/v			
Seminal Fluid	Seminal Fluid	5.0% v/v			
Mucus	Mucin	0.8% w/v			



Class/Substance	Active Ingredient	Concentration Tested		
	Benzocaine 5%; Resorcinol 2%	0.25% w/v		
	Clotrimazole 2%	0.25% w/v		
	Miconazole Nitrate 2%	0.25% w/v		
	Tioconazole	0.25% w/v		
	5% w/w Aciclovir	0.25% w/v		
Over the counter(OTC) Vaginal Products; Contraceptives; Vaginal treatments	Glycerin, Propylene glycol	0.25% w/v		
	Glycerin; Carbomer	0.12% w/v		
	Glycerin, Hydroxyethyl cellulose	0.25% w/v		
	Goldenseal 3X HPUS; Kreosotum 12X HPUS	0.25% w/v		
	Povidone-iodine 10%	0.25% v/v		
	Nonoxynol-9 12.5%	0.25% w/v		
lemorrhoidal Cream	Glycerin 14%; Pramoxine HCl 1%	0.25% w/v		
eukocytes	Leukocytes	10 ⁵ cells/mL		
ntravaginal Hormones	Progesterone; Estradiol	7 mg/mL Progesterone + 0.07 mg/ mL Beta Estradiol		

a. In tests with substances diluted into pooled T. vaginalis-positive swab matrix, test interference was observed in tests with blood at 60% v/v. No test interference was observed in tests with blood at 50% v/v. This is addressed in Limitations of the Procedure, Limitations.

Carry-Over Contamination Study

This study was conducted to demonstrate that single-use, self-contained GeneXpert cartridges prevent carryover contamination in negative samples run after very high positive samples in the same GeneXpert module. A negative sample (*T. vaginalis* negative vaginal swabs in Cepheid Xpert Swab Transport Reagent) was run followed by 20 rounds of high positive sample (*T. vaginalis* ATCC 30001 at 10⁶ cells/mL diluted in vaginal swab matrix) alternating with a negative sample in two separate GeneXpert modules for a total of 40 high positive and 42 negative samples for each module. This testing scheme resulted in a total of 82 runs (40 positive + 42 negative samples). There was no evidence of carry-over contamination as all 40 positive samples were correctly reported as **TV DETECTED** and all 42 negative samples were correctly reported as **TV NOT DETECTED**.

Reproducibility

Intra-site reproducibility of the Xpert TV Test was evaluated at three sites (two external, one in-house). Site 1 used an Infinity-80 instrument. Sites 2 and 3 used GeneXpert Dx instruments. Specimens were created by spiking *Trichomonas vaginalis* (ATCC 30001) into pooled, *Trichomonas vaginalis* negative urine (patient urine added to Cepheid Urine Transport Reagent) or vaginal swab matrix (vaginal swabs collected into Cepheid Swab Transport Reagent). The specimens were prepared at concentration levels representing high negative (below LoD), LoD (~1X LoD), moderate positive (~3X LoD), and negative (*Trichomonas vaginalis* negative clinical matrix). A panel of 8 specimens (4 in urine and 4 in vaginal swab matrix) was tested twice per day, on 12 different days, by two different operators, at each of three sites (8 specimens x 2 replicates x 12 days x 2 operators x 3 sites = 1,152 observations total). Three lots of Xpert TV Test cartridges were used at each of the 3 testing sites, with each lot used for 4 days of testing. Positive and negative controls were included in the study. The Xpert TV Test was performed according to the Xpert TV Test procedure. The rate of agreement with expected results is shown by site in Table 14.



6 1 3	Site 1 (Infinity-80)			Site 2	Site 2 (GeneXpert Dx)			(GeneXpe	Total Agreement		
Sample ^a	Op 1	Op 2	Site	Op 1	Op 2	Site	Op 1	Op 2	Site	by Sample	
FS-Neg	100% (24/24)	100% (24/24)	100% (48/48)	100% (24/24)	100% (24/24)	100% (48/48)	100% (24/24)	100% (24/24)	100% (48/48)	100% (144/144)	
FS-Mod Pos (~3X LoD; ~6	100%	100%	100%	100%	100%	100%	100%	100%	100%	100% (144/144)	
cells/mL)	(24/24)	(24/24)	(48/48)	(24/24)	(24/24)	(48/48)	(24/24)	(24/24)	(48/48)		
FS-LoD (~1X LoD; ~2 cells/	95.8%	100%	97.9%	87.5%	95.8%	91.7%	100%	95.8%	97.9%	95.8% (138/144)	
mL)	(23/24)	(24/24)	(47/48)	(21/24)	(23/24)	(44/48)	(24/24)	(23/24)	(47/48)		
FS-High Neg (below LoD;	87.5%	75.0%	81.3%	66.7%	79.2%	72.9%	79.2%	70.8%	75.0%	76.4% (110/144)	
<2 cells/mL)	(21/24)	(18/24)	(39/48)	(16/24)	(19/24)	(35/48)	(19/24)	(17/24)	(36/48)		
UR-Neg	100% (24/24)	100% (24/24)	100% (48/48)	100% (24/24)	100% (24/24)	100% (48/48)	100% (24/24)	100% (24/24)	100% (48/48)	100% (144/144)	
UR-Mod Pos (~3X LoD; ~9	100%	100%	100%	100%	100%	100%	100%	100%	100%	100% (144/144)	
cells/mL)	(24/24)	(24/24)	(48/48)	(24/24)	(24/24)	(48/48)	(24/24)	(24/24)	(48/48)		
UR-LoD (~1X LoD; ~3	75.0%	91.7%	83.3%	83.3%	91.3%	87.2%	91.7%	100%	95.8%	88.8% (127/143)	
cells/mL)	(18/24)	(22/24)	(40/48)	(20/24)	(21/23) ^b	(41/47)	(22/24)	(24/24)	(46/48)		
UR-High Neg (below LoD;	75.0%	75.0%	75.0%	70.8%	54.2%	62.5%	75.0%	75.0%	75.0%	70.8% (102/144)	
< 3 cells/mL)	(18/24)	(18/24)	(36/48)	(17/24)	(13/24)	(30/48)	(18/24)	(18/24)	(36/48)		

Table 14. Summary of Reproducibility Results

a. FS=female swab matrix; UR=urine matrix.

b. One sample indeterminate on initial and retest.

The reproducibility of the Xpert TV Test was also evaluated in terms of the fluorescence signal expressed in Ct values for each target detected. The mean, standard deviation (SD), and coefficient of variation (CV) between-sites, between-lots, between-days, between-operators, and residual variability for each panel member are presented in Table 15.



Sampleª	Test Channel (Analyte) N	N ^b	Mean	Between- Site		Between- Lot		Between- Day		Between- Operator		Residual		Total	
		IN ²	Ct	SD	CV (%) ^c	SD	CV (%) ^c	SD	CV (%) ^c	SD	CV (%) ^د	SD	CV (%) ^c	SD	CV (%) ^c
FS-Neg	SPC	144	33.7	0.0	0.0	0.1	23.2	0.1	8.9	0.0	0.0	0.4	67.9	0.4	1.2
FS-Mod Pos (~3X LoD; ~6 cells/mL)	TV	144	35.4	0.1	7.9	0.0	0.0	0.0	0.0	0.1	12.5	0.8	79.7	0.8	2.3
FS-LoD (~1X LoD; ~ 2 cells/ mL)	TV	138	38.5	0.0	0.0	0.0	0.0	0.5	28.0	0.0	0.0	1.2	72.0	1.3	3.5
FS-High Neg (below LoD; < 2 cells/mL)	TV	110	39.4	0.0	0.0	0.0	0.0	0.4	17.6	0.0	0.0	1.7	82.4	1.8	4.5
UR-Neg	SPC	144	33.9	0.1	8.6	0.0	0.0	0.1	9.0	0.1	18.5	0.4	63.9	0.4	1.2
UR-Mod Pos (~3X LoD; ~9 cells/mL)	TV	144	35.5	0.2	22.3	0.1	9.6	0.0	0.0	0.0	0.0	0.6	67.9	0.7	1.9
UR-LoD (~1X LoD; ~3 cells/ mL)	TV	127	39.3	0.0	0.0	0.4	24.4	0.0	0.0	0.0	0.0	1.2	75.6	1.3	3.4
UR-High Neg (below LoD; < 3 cells/mL)	TV	102	39.0	0.0	0.0	0.3	14.4	0.7	29.5	0.3	11.6	1.0	44.6	1.3	3.3

Table 15. Summary of Reproducibility Data

a. FS=female swab matrix; UR=urine matrix.

b. Results with non-zero Ct values out of 144.

c. (%) is contribution of variance component to overall CV.

Instrument System Precision

An in-house precision study was conducted to compare the performance of the GeneXpert Dx and the GeneXpert Infinity Instrument Systems using specimens comprised of *Trichomonas vaginalis* (ATCC[®]

30001[™]) spiked into negative urine (patient urine added to Cepheid Urine Transport Reagent) or vaginal swab matrix (vaginal swabs collected into Cepheid Swab Transport Reagent). The specimens were prepared at concentration levels representing high negative (below LoD), LoD (~1X LoD), moderate positive (~3X LoD), and negative (*Trichomonas vaginalis* negative clinical matrix). A panel of 8 specimens (4 in urine matrix and 4 in vaginal swab matrix) was tested on 12 different days by two operators. Each operator conducted four runs of each panel specimen per day on each of the three instrument systems (8 specimens x 4 times/day x 12 days x 2 operators x 3 instrument systems = 2,304 observations total). Three lots of Xpert TV Test cartridges were used for the study, with each lot used for 4 days of testing. Positive and negative controls were included in the study. The Xpert TV Test was performed according to the Xpert TV Test procedure. The rate of agreement with expected results is shown by instrument in Table 16.

c 1 a	GeneXpert Dx				Infinity-48	;		Infinity-80	% Total Agreement	
Sample ^a	Op 1	Op 2	Inst	Op 1	Op 2	Inst	Op 1	Op 2	Inst	by Sample
FS-Neg	100% (48/48)	100% (48/48)	100% (96/96)	97.9% (47/48)	100% (48/48)	99.0% (95/96)	100% (48/48)	100% (48/48)	100% (96/96)	99.7% (287/288)
FS-Mod Pos (~3X LoD; ~6 cells/mL)	100% (48/48)	100% (48/48)	100% (96/96)	100% (48/48)	100% (48/48)	100% (96/96)	100% (48/48)	100% (48/48)	100% (96/96)	100% (288/288)
FS-LoD (~1X LoD; ~ 2 cells/mL)	93.8% (45/48)	87.5% (42/48)	90.6% (87/96)	93.8% (45/48)	89.6% (43/48)	91.7% (88/96)	95.8% (46/48)	89.6% (43/48)	92.7% (89/96)	91.7% (264/288)

Table 16.	Summary	∕ of Pre	cision	Results
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Sample ^a	GeneXpert Dx				Infinity-48	}		Infinity-80	% Total Agreement	
	Op 1	Op 2	Inst	Op 1	Op 2	Inst	Op 1	Op 2	Inst	by Sample
FS-High Neg (below LoD;	74.5%	75.0%	74.7%	77.1%	75.0%	76.0%	83.3%	68.8%	76.0%	75.6% (217/287) ^b
< 2 cells/mL)	(35/47)	(36/48)	(71/95)	(37/48)	(36/48)	(73/96)	(40/48)	(33/48)	(73/96)	
UR-Neg	100% (48/48)	100% (48/48)	100% (96/96)	100% (48/48)	100% (48/48)	100% (96/96)	100% (48/48)	100% (47/47)	100% (95/95)	100% (287/287) ^b
UR-Mod Pos (~3X LoD; ~9	100%	100%	100%	100%	100%	100%	100%	100%	100%	100% (288/288)
cells/mL)	(48/48)	(48/48)	(96/96)	(48/48)	(48/48)	(96/96)	(48/48)	(48/48)	(96/96)	
UR-LoD (~1X LoD; ~3	93.8%	93.8%	93.8%	95.8%	89.6%	92.7%	95.8%	95.8%	95.8%	94.1% (271/288)
cells/mL)	(45/48)	(45/48)	(90/96)	(46/48)	(43/48)	(89/96)	(46/48)	(46/48)	(92/96)	
UR-High Neg (below	72.9%	77.1%	75.0%	70.8%	79.2%	75.0%	81.3%	85.4%	83.3%	77.8% (224/288)
LoD; < 3 cells/mL)	(35/48)	(37/48)	72/96)	(34/48)	(38/48)	(72/96)	(39/48)	(41/48)	(80/96)	

a. FS=female swab matrix; UR= urine matrix.

b. One FS-Low-Pos and one UR-Neg sample indeterminate and not retested.

The precision of the Xpert TV Test was also evaluated in terms of the fluorescence signal expressed in Ct values for each target detected. The mean, standard deviation (SD), and coefficient of variation (CV) between-instruments, between-lots, between-days, between-operators, and residual variability for each panel member are presented in Table 17.

Sampleª	Test Channel (Analyte)	N Þ	Mean Ct	Between- Instrument		Between- Lot		Between- Day		Between- Operator		Residual		Total	
				SD	CV (%) ^c	SD	CV (%) ^c	SD	CV (%) ^c	SD	CV (%) ^c	SD	CV (%) ^c	SD	CV (%) ^c
FS-Neg	SPC	288	31.9	0.0	0.0	0.3	53.5	0.0	0.0	0.1	1.9	0.2	44.6	0.4	1.1
FS-Mod Pos (~3X LoD; ~6 cells/mL)	τv	288	35.2	0.0	0.0	0.3	22.4	0.0	0.0	0.1	4.5	0.4	73.1	0.5	1.5
FS-LoD (~1X LoD; ~2 cells/mL)	τv	264	39.0	0.2	3.3	0.1	0.4	0.2	1.3	0.0	0.0	1.3	95.0	1.3	3.4
FS-High Neg (below LoD; < 2 cells/mL)	τv	217	39.4	0.0	0.0	0.0	0.0	0.0	0.0	0.2	1.6	1.3	98.4	1.3	3.2
UR-Neg	SPC	287	32.4	0.0	0.0	0.3	47.2	0.1	2.9	0.0	0.0	0.3	49.9	0.4	1.2
UR-Mod Pos (~3X LoD; ~9 cells/mL)	τv	288	35.4	0.0	0.0	0.4	30.4	0.0	0.0	0.2	11.3	0.5	58.3	0.6	1.8
UR-LoD (~1X LoD; ~3 cells/mL)	τv	271	38.2	0.0	0.0	0.5	13.6	0.6	16.2	0.3	3.6	1.2	66.5	1.4	3.7
UR-High Neg (below LoD; < 3 cells/mL)	τv	224	38.9	0.0	0.0.	0.3	5.4	0.0	0.0	0.3	4.2	1.2	90.3	1.3	3.3

Table 17. Summary of Precision Data

a. FS=female swab matrix; UR=urine matrix

b. Results with non-zero Ct values out of 288.

c. (%) is contribution of variance component to overall CV.

? Appendix

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Table of Symbols

Symbol	Meaning
R _{konly}	For prescription use only
REF	Catalog number
IVD	In vitro diagnostic medical device
8	Do not reuse
LOT	Batch code
i	Consult instructions for use
<u>^</u>	Caution
	Manufacturer
53	Country of manufacture
Σ	Contains sufficient for <i>n</i> tests
CONTROL	Control
	Expiration date
X	Temperature limitation
620	Biological risks
$\langle \mathbf{\hat{k}} \rangle$	Warning



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Revision History

Description of Changes: 303-9043, Rev. A

Purpose: Initial release