# AmpliSens® T.vaginalis / N.gonorrhoeae-**MULTIPRIME-FRT** PCR kit



For Professional Use Only

# Instruction Manual

#### **KEY TO SYMBOLS USED**

REF	Catalogue number	$\triangle$	Caution
LOT	Batch code	Σ	Sufficient for
IVD	In vitro diagnostic medical device	><	Use-by Date
VER	Version	$\bigcap$ i	Consult instructions for use
$\bigwedge$	Temperature limit	**	Keep away from sunlight
***	Manufacturer	NCA	Negative control of amplification
$\sim$	Date of manufacture	C-	Negative control of extraction
EC REP	Authorized representative in the European Community	C+	Positive control of amplification
		IC	Internal control

#### 1. INTENDED USE

AmpliSens® T.vaginalis / N.gonorrhoeae-MULTIPRIME-FRT PCR kit is an in vitro nucleic acid amplification test for simultaneous detection of *Trichomonas vaginalis* and *Neisseria* gonorrhoeae DNA in the clinical material (urogenital, rectal, and oropharyngeal swabs; conjunctival discharge; urine samples, prostate gland secretion) using real-time hybridization-fluorescence detection of amplified products.

The results of PCR analysis are taken into account in complex diagnostics of NOTE:

#### 2. PRINCIPLE OF PCR DETECTION

 ${\it T.vaginalis / N.gonorrhoeae} \ {\it detection} \ {\it by} \ {\it the polymerase chain reaction} \ ({\it PCR}) \ is \ {\it based on} \ the \ amplification \ of \ the \ pathogen \ genome \ specific \ region \ using \ specific \ {\it T.vaginalis / N.gonorrhoeae} \ {\it the pathogen } \ {\it the path$ N gonorrhoeae primers. In the real-time PCR, the amplified product is detected with the use of fluorescent dyes. These dyes are linked to oligonucleotide probes which bind specifically to the amplified product during thermocycling. The real-time monitoring of fluorescence intensities during the real-time PCR allows the detection of accumulating product without re-

opening the reaction tubes after the PCR run.

AmpliSens® T.vaginalis / N.gonorrhoeae-MULTIPRIME-FRT PCR kit is a qualitative test that contains the Internal Control (Internal Control-FL (IC)). It must be used in the extraction procedure in order to control the extraction process of each individual sample and to identify possible reaction inhibition.

possible reaction inhibition.

AmpliSens® T.vaginalis / N.gonorrhoeae-MULTIPRIME-FRT PCR kit uses "hot-start," which greatly reduces the frequency of nonspecifically primed reactions. "Hot-start" is guaranteed by using chemically modified polymerase (TaqF). The chemically modified polymerase (TaqF) is activated by heating at 95 °C for 15 min.

The PCR kit contains the system for prevention of contamination by amplicons using the enzyme uracil-DNA-glycosylase (UDG) and dUTP. The enzyme UDG recognizes and catalyzes the destruction of the DNA containing deoxyuridine, but has no effect on DNA containing deoxythymidine. Deoxyuridine is absent in the authentic DNA, but is always present in amplicons, because dUTP is a part of dNTP mixture in the reagents for the amplification. Due to the deoxyuridine containing contaminating amplicons are sensitive to the destruction by UDG before the DNA-target amplification. So the amplicons cannot be amplified.

The enzyme UDG is thermolabile. It is inactivated by heating at temperature above 50 °C. Therefore, UDG does not destroy the target amplicons which are accumulated during PCR.

Table 1

Channel for fluorophore	FAM	JOE	ROX
DNA-target	Trichomonas vaginalis	Neisseria gonorrhoeae	Internal Control- FL (IC) DNA
Target gene	Trichomonas vaginalis repeated DNA target for PCR identification	gene 16S rRNA	Artificially synthesized sequence

#### 3. CONTENT

AmpliSens® T.vaginalis / N.gonorrhoeae-MULTIPRIME-FRT PCR kit is produced in

variant FRT-100 F, REF R-B65-F(RG,iQ)-CE.

Reagent	Description	Volume, ml	Quantity
PCR-mix-1-FL T.vaginalis / N.gonorrhoeae	clear liquid from colorless to light lilac colour	1.2	1 tube
PCR-mix-2-FRT	colorless clear liquid	0.3	2 tubes
Polymerase (TaqF)	colorless clear liquid	0.03	2 tube
Positive Control complex (C+)	colorless clear liquid	0.2	1 tube
DNA-buffer	colorless clear liquid	0.5	1 tube
Negative Control (C-)*	colorless clear liquid	1.2	1 tube
Internal Control-FL (IC)**	colorless clear liquid	1.0	1 tube

- must be used in the extraction procedure as Negative Control of Extraction. add 10 µl of Internal Control-FL (IC) during the DNA extraction procedure directly to the sample/lysis mixture (see the DNA-sorb-AM protocol). Variant FRT-100 F is intended for 110 reactions (including controls)

#### 4. ADDITIONAL REQUIREMENTS

- DNA extraction kit.
- Transport medium
- Disposable powder-free gloves and laboratory coat.
- Pipettes (adjustable).
- Sterile pipette tips with aerosol filters (up to 100 µl).
- Tube racks.
- Vortex mixer
- Desktop centrifuge with a rotor for 2 ml reaction tubes.
- PCR box.
- Real-time instruments (for example, Rotor-Gene 3000/6000 (Corbett Research, Australia); Rotor-Gene Q (QIAGEN, Germany); iCycler iQ5 (Bio-Rad, USA); Mx3000P (Stratagene, USA).
- Disposable polypropylene tubes:
- a) thin-walled 0.2-ml PCR tubes with domed caps if a plate-type instrument is used; b) thin-walled 0.2-ml PCR tubes with flat caps or strips of four 0.1-ml Rotor-Gene PCR
- tubes if a rotor-type instrument is used Refrigerator for 2–8 °C.
- Deep-freezer at the temperature from minus 24 to minus 16 °C.
- Reservoir for used tips.

#### 5. GENERAL PRECAUTIONS

The user should always pay attention to the following:

- Use sterile pipette tips with aerosol filters and use a new tip for every procedure. Store all extracted positive material (specimens, controls and amplicons) away from all
- other reagents and add it to the reaction mix in a distantly separated facility. Thaw all components thoroughly at room temperature before starting an assay.
- When thawed, mix the components and centrifuge briefly.

  Use disposable protective gloves and laboratory cloths, and protect eyes while samples and reagents handling. Thoroughly wash hands afterwards.
- Do not eat, drink, smoke, apply cosmetics, or handle contact lenses in laboratory work
- Do not use a kit after its expiration date.
- Dispose of all specimens and unused reagents in accordance with local regulations.
- Samples should be considered potentially infectious and handled in biological cabinet in compliance with appropriate biosafety practices.
- Clean and disinfect all samples or reagents spills using a disinfectant, such as 0.5 % sodium hypochlorite or another suitable disinfectant.
- Avoid inhalation of vapors, samples and reagents contact with the skin, eyes, and mucous membranes. Harmful if swallowed. If these solutions come into contact, rinse the injured area immediately with water and seek medical advice if necessary. Safety Data Sheets (SDS) are available on request.
- Use of this product should be limited to personnel trained in DNA amplification techniques.
- Workflow in the laboratory must be one-directional, beginning in the Extraction Area and moving to the Amplification and Detection Area. Do not return samples, equipment and reagents in the area where the previous step was performed.



Some components of this kit contain sodium azide as a preservative. Do not use metal tubing for reagent transfer.

#### 6. SAMPLING AND HANDLING

Obtaining samples of biological materials for PCR-analysis, transportation and storage are described in the manufacturer's handbook [1]. It is recommended that this handbook is read before starting work.

AmpliSens® T.vaginalis / N.gonorrhoeae-MULTIPRIME-FRT PCR kit is intended for analysis of the DNA extracted with DNA extraction kits from the clinical material (urogenital, rectal, and oropharyngeal swabs; conjunctival discharge; urine samples (a sediment of the first portion of the morning specimen); prostate gland secretion).

#### 7. WORKING CONDITIONS

AmpliSens® T.vaginalis / N.gonorrhoeae-MULTIPRIME-FRT PCR kit should be used at

## 8. PROTOCOL

#### 8.1. DNA Extraction

It is recommended to use the following nucleic acid extraction kits:

- DNA-sorb-AM:
- For other nucleic acid extraction kits see Guidelines [2]

The DNA extraction of each test sample is carried out in the presence of Internal Control-FL (IC)

extraction procedure it is necessary to carry out the control reactions as follows:

- Add 100 µl of Negative Control (C-) to the tube labeled C- (Negative C-

NOTE: Extract DNA according to the manufacturer's protocol

#### 8.2. Preparing PCR

The type of tubes depends on the PCR instrument used for analysis.
Use disposable filter tips for adding reagents, DNA and control samples into tubes.

#### Preparing tubes for PCR

The total reaction volume is 25 µI, the volume of DNA sample is 10 µI.

- Thaw the tube with PCR-mix-2-FRT. Vortex the tubes with PCR-mix-1-FL *T.vaginalis* / *N.gonorrhoeae*, PCR-mix-2-FRT, and polymerase (TaqF) and then centrifuge briefly. Take the required number of tubes/strips for amplification of the DNA obtained from iake the required number of tubes/strips for amplification clinical and control samples. For N reactions (including 2 controls) add to a new tube: 10-(N+1) µl of PCR-mix-1-FL *T.vaginalis / N.gonorrhoeae*, 5.0-(N+1) µl of PCR-mix-2-FRT,

0.5:(N+1) µl of polymerase (TaqF).

Vortex the tube, then centrifuge it briefly. Transfer 15 µl of the prepared mixture to each

- Using tips with aerosol filter, add  $10~\mu l$  of DNA samples obtained at the DNA extraction stage
- stage. Carry out the control amplification reactions: NCA Add 10 µl of DNA-buffer to the tube labeled NCA (Negative Control of NCA
- Add 10 ul of Positive Control complex to the tube labeled C+ (Positive C+ control of Amplification).
- C-Add 10 µl of the sample extracted from the Negative Control (C-)

reagent to the tube labeled C- (Negative control of Extraction

#### 8.2.2. Amplification

1. Create a temperature profile on your instrument as follows:

Table 2

Amprisens-1 amplification program						
	Rotor-type Instruments <sup>1</sup>		Plate-type Instruments <sup>2</sup>			
Step	Temperature, °C	Time	Cycles	Temperature, °C	Time	Cycles
Hold	95	15 min	1	95	15 min	1
	95	5 s		95	5 s	
Cycling	60	20 s	5	60	20 s	5
	72	15 s		72	15 s	
	95	5 s		95	5 s	
Cycling 2	60	20 s Fluorescence acquiring	40	60	30 s Fluorescence acquiring	40
	72	15 s	l	72	15 s	

AmpliSens-1 amplification program

Fluorescent signal is detected in the channels designed for the FAM. JOE and ROX fluorophores (other channels are enabled if several tests are simultaneously carried out in a single run).

- 2. Adjust the fluorescence channel sensitivity according to the *Important Product Information Bulletin* and Guidelines [2].

  3. Insert tubes into the reaction module of the device.

  4. Run the amplification program with fluorescence detection.

- Analyze results after the amplification program is completed.

#### 9. DATA ANALYSIS

Analysis of results is performed by the software of the real-time PCR instrument used by measuring fluorescence signal accumulation in three channels:

- The signal of the *Trichomonas vaginalis* DNA amplification product is detected in the channel for the FAM fluorophore.
- The signal of the Neisseria gonorrhoeae DNA amplification product is detected in the channel for the JOE fluorophore.
- The signal of the IC DNA amplification product is detected in the channel for the ROX

Results are interpreted by the crossing (or not-crossing) the fluorescence curve with the threshold line set at the specific level that corresponds to the presence (or absence) of a Ct value of the DNA sample in the corresponding column of the results grid

- value of the DNA sample in the corresponding column of the results grid.
  Principle of interpretation is the following:
  Trichomonas vaginalis DNA is detected if the Ct value is determined in the results grid in the channel for the FAM fluorophore. Moreover, the fluorescence curve of the sample should cross the threshold line in the area of typical exponential growth of fluorescence.
  Neisseria gonorrhoeae DNA is detected if the Ct value is determined in the results grid in the channel for the JOE fluorophore. Moreover, the fluorescence curve of the sample should cross the threshold line in the area of typical exponential growth of fluorescence.
  Trichomonas vaginalis and Neisseria gonorrhoeae DNA are not detected in a sample if the Ct value is not determined (absent) in the channels for FAM and JOE fluorophores, whereas the Ct value determined in the channel for the ROX fluorophore is less than the boundary Ct value specified in the Important Product Information Bulletin less than the boundary Ct value specified in the Important Product Information Bulletin
- The result is invalid if the Ct value is not determined (absent) in the channels for FAM, JOE or ROX fluorophores. In such cases, the PCR analysis should be repeated.

Boundary Ct values are specified in the Important Product Information Bulletin enclosed to the PCR kit. See also Guidelines [2].

The result of the analysis is considered reliable only if the results obtained for the Positive and Negative Controls of amplification as well as for the Negative Control of extraction are correct (see Table 3).

Table 3

Control	011	Ct value in the channel for fluorophore		
Control	Stage for control	FAM, JOE	ROX	
c-	DNA extraction	Absent	< boundary value	
NCA	PCR	Absent	Absent	
C+	PCR	< boundary value	< boundary value	

#### 10. TROUBLESHOOTING

Results of analysis are not taken into account in the following cases:

- If the Ct value determined for the Positive Control of Amplification (C+) in the channels for the FAM and\or JOE and\or ROX fluorophores is greater than the boundary Ct value
- or the FAM and/or JOE and/or ROX fluorophores is greater than the boundary Cr value or absent, the amplification should be repeated for all samples in which the boundary Cr is absent in the channels for the FAM and/or JOE and/or ROX fluorophores.

  2. If the Cr value is determined for the Negative Control of Amplification (NCA) and/or Negative Control of Extraction (C-) in the channels for the FAM and/or JOE fluorophores, the PCR analysis should be repeated for all samples in which the Cr value is determined in the channels for the FAM and/or JOE fluorophores.

  If you have any further questions or if encounter problems, please contact our Authorized

representative in the European Community.

#### 11. TRANSPORTATION

AmpliSens® T.vaginalis / N.gonorrhoeae-MULTIPRIME-FRT PCR kit should be transported at 2–8 °C for no longer than 5 days

#### 12. STABILITY AND STORAGE

All components of the AmpliSens® *T.vaginalis / N.gonorrhoeae*-MULTIPRIME-FRT PCR kit are to be stored at 2–8 °C when not in use (except for polymerase (TaqF) and PCR-mix-

All components of the AmpliSens® *T.vaginalis / N.gonorrhoeae*-MULTIPRIME-FRT PCR kit are stable until the expiry date on the label. PCR kit variant FRT-100 F can be stored without unpacking at 2 to 8 °C for 3 months from the date of manufacture before opening. Once opened, PCR kit variant FRT-100 F should be unpacked in accordance with the storage temperatures for each component. The shelf life of reagents before and after the first use is the same, unless otherwise stated.

Polymerase (TaqF) and PCR-mix-2-FRT are to be stored at the temperature from minus 24 to minus 16  $^{\circ}\text{C}$  when not in use. NOTE:

NOTE: PCR-mix-1-FL T.vaginalis / N.gonorrhoeae is to be kept away from light.

#### 13. SPECIFICATIONS

## 13.1. Sensitivity

The analytical sensitivity of AmpliSens® T.vaginalis / N.gonorrhoeae-MULTIPRIME-FRT PCR kit is the following:

Clinical material	Transport medium	Nucleic acid extraction kit	Microorganism	Sensitivity, GE/ml <sup>3</sup>
Urogenital	Transport medium for swabs or Transport medium with	DNA-sorb-AM	Trichomonas vaginalis	5x10 <sup>2</sup>
swabs	mucolytic agent		Neisseria gonorrhoeae	5x10 <sup>2</sup>
Urine <sup>4</sup>	_	DNA-sorb-AM	Trichomonas vaginalis	10 <sup>3</sup>
Offine			Neisseria gonorrhoeae	10 <sup>3</sup>

The analytical sensitivity of each microorganism does not change even at high concentrations of the other microorganism (to 10° GE/ml). NOTE:

<sup>2</sup> For example, iQ5, Mx3000P, Mx3000

<sup>3</sup> The quantity of genome equivalents of microorganism per 1 ml of the sample.

For example, Rotor-Gene 3000, Rotor-Gene 6000, Rotor-Gene Q.

#### 13.2. Specificity

The analytical specificity of AmpliSens® T.vaginalis / N.gonorrhoeae-MULTIPRIME-FRT PCR kit is ensured by the selection of specific primers and probes as well as strict reaction conditions. The primers and probes have been checked for possible homologies to all sequences published in gene banks by sequence comparison analysis.

Nonspecific responses were absent while testing human DNA samples as well as a DNA condition of the following princeposings: Organization variables in the following princeposings:

Nonspecific responses were absent while testing human DNA samples as well as a DNA panel of the following microorganisms: Gardnerella vaginalis, Lactobacillus spp., Escherichia coli, Staphylococcus spp., Streptococcus spp., Candida albicans, Mycoplasma hominis, Ureaplasma urealyticum, Ureaplasma parvum, Mycoplasma genitalium, Chlamydia trachomatis, Neisseria spp., Neisseria gonorrhoeae, Trichomonas vaginalis, Treponema pallidum, Toxoplasma gondii, HSV types 1 and 2, CMV, and HPV.

The clinical specificity of AmpliSens® T.vaginalis / N.gonorrhoeae-MULTIPRIME-FRT PCR kit was confirmed in laboratory clinical trials.

#### 14. REFERENCES

- Handbook "Sampling, Transportation, and Storage of Clinical Material for PCR Diagnostics", developed by Federal Budget Institute of Science "Central Research Institute for Epidemiology" of Federal Service for Surveillance on Consumers' Rights Protection and Human Well-Being.
- Protection and Human well-being.

  Guidelines "Real-Time PCR Detection of STIs and Other Reproductive Tract Infections", developed by Federal Budget Institute of Science "Central Research Institute for Epidemiology" of Federal Service for Surveillance on Consumers' Rights Protection and Human Well-Being.

## **15. QUALITY CONTROL**

In compliance with Federal Budget Institute of Science "Central Research Institute for Epidemiology" ISO 13485-Certified Quality Management System, each lot of **AmpliSens® T.vaginalis / N.gonorrhoeae-MULTIPRIME-FRT** PCR kit has been tested against predetermined specifications to ensure consistent product quality.

List of Changes Made in the Instruction Manual

VER	Location of changes	Essence of changes		
29.06.11 LA	Cover page, text	The name of Institution was changed to Federal Budget Institution of Science "Central Research Institute for Epidemiology"		
	Through the text	Corrections in accordance with the template		
	1. Intended use	The clinical material was specified		
17.02.16 PM	8.1. DNA extraction	Information about controls of extraction was added		
FIVI	9. Data analysis	The sections were rewritten		
	<ol><li>Troubleshooting</li></ol>	The sections were rewritten		
15.03.18 PM	Footer, 3. Content	REF R-B65(iQ)-CE was deleted		
10.09.18 EM	3. Content	The colour of the reagent was specified		
17.01.19 PM	Principle of PCR detection	The information about the enzyme UDG was added. The information about "hot-start" was corrected		
	Through the text	The text formatting was changed		
29.04.20 KK	Principle of PCR detection	The table with targets was added.		
NN.	Footer	The phrase "Not for use in the Russian Federation" was added		
30.10.20 MA	Through the text, Footer	The information about variant FRT REF R-B65(RG)-CE was deleted		
18.03.21 MM		The name, address and contact information for Authorized representative in the European Community was changed		
30.11.21 MM	12. Stability and storage	The information about storage conditions for 3 months from the date of manufacture and subsequent unpacking was added		
	Through the text	The reference numbers of nucleic acid extraction kits and transport mediums were deleted		

## AmpliSens®



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