



# STD Direct Flow Chip Kit

# Detection of pathogens responsible for sexually transmitted diseases (STD) through multiplex PCR and reverse hybridization

For all hybriSpot platforms

<u>Compatible with version 2.2.0 of hybriSoft HSHS and later and lyophilized format of the kit.</u> For compatibility with other versions, please contact the manufacturer / supplier.



Ref. MAD-003938M-HS12 Ref. MAD-003938M-HS Σ

24 determinations 24 determinations

For in vitro diagnostic use only\* Directive 98/79/EC

\*The 0318 Notified Organism only intervenes in the evaluation of the compliance of the test for Chlamydia trachomatis Biovars A-K in urine and semen samples; urethral, endocervical, vaginal and anal swabs; the rest of pathogens have the self-certified CE marking.





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# **1. INTENDED USE**

STD Direct Flow Chip is an in vitro diagnostic kit for the detection of pathogenic organisms causing sexually transmitted diseases (STD) in humans. The organisms causing these infections are often difficult to detect and include viruses, bacteria or parasites, usually occurring co-infections. The methods that are currently used for their diagnosis are very laborious and not always show a 100% specificity. The STD Direct Flow Chip allows the simultaneous detection of 11 pathogens: Chlamydia trachomatis<sup>1\*</sup>, Haemophilus ducreyi, Herpes simplex virus 1 and Herpes simplex virus 2, Mycoplasma genitalium, Mycoplasma hominis, Treponema pallidum, Trichomonas vaqinalis Neisseria gonorrhoeae, and Ureaplasma<sup>∠</sup> (urealyticum/parvum). The STD Direct Flow Chip kit allows the direct detection of these infectious agents from different types of clinical samples (urine, semen, liquid-based endocervical cytologies; and urethral, endocervical, vaginal, anal and pharyngeal swabs) without previous extraction of DNA, as well as from purified DNA of these type of clinical samples.

<sup>1</sup>Biovars and Serovars of the *Chlamydia trachomatis* species detected with the *STD Direct Flow Chip* kit:

- Chlamydia trachomatis Biovar Trachoma: Serovars A-K.
- Chamydia trachomatis Biovar LGV: Serovars L1-L3.

\* The 0318 Notified Organism only intervenes in the evaluation of the compliance of the test for Chlamydia trachomatis Biovars A-K in urine samples; urethral, endocervical and anal swabs. The detection of Chlamydia trachomatis Biovars L1-L3, as well as the detection of the Biovars A-K in seminal and pharyngeal samples are not regulated by the certification of the 0318 Notified Organism.

<sup>2</sup>Ureaplasmas detected with the *STD Direct Flow Chip* kit:

- Ureaplasma urealyticum: UUR2, UUR4, UUR5, UUR7, UUR8, UUR9, UUR10, UUR11, UUR12, UUR13.
- Ureaplasma parvum: UPA1, UPA3, UPA6, UPA14.

Note: In the literature, it is common to use the terms Biovar or Serovar indistinctly to refer to Serovars A-K of the Trachoma Biovar and L1-L3 of the Biovar LGV of the *C. trachomatis* species.

Microbiological status: Product not sterile

# **2. PRINCIPLE OF THE METHOD**

The *STD Direct Flow Chip* kit is based on a methodology consisting of the simultaneous amplification of bacterial, viral and protozoal DNA by multiplex PCR in only one step and directly from cellular extracts, followed by hybridization on a membrane with specific DNA probes using the DNA-Flow technology for HybriSpot platforms, both automated and manual. The biotinylated amplicons generated after the multiplex PCR are hybridized in membranes containing an array of specific probes for each human pathogen, as well as amplification and hybridization control probes. Unlike conventional hybridization on surfaces, the *DNA-Flow* technology allows a very fast binding of the PCR product and its specific probe in a three-dimensional porous environment. Once the binding has occurred between the specific amplicons and their corresponding probes, the signal is visualized by a colorimetric immunoenzymatic reaction with Streptavidin-Phosphatase and a chromogen (NBT-BCIP), which generates insoluble precipitates on the membrane in the positions in which there has been hybridization. The results are analyzed automatically







with the HybriSoft software.

# **3. COMPONENTS**

The **STD Direct Flow Chip** kit is marketed in two main formats according to the type of hybridization platform to be used for the analysis of clinical samples. Both formats provide the necessary reagents for the amplification through multiplex PCR and subsequent hybridization of 24 clinical samples. Each kit format contains the following components and references:

KIT/COMPONENTS	FORMAT	REFERENCES
STD Direct Flow Chip kit (Manual)	24 tests	MAD-003938M-HS12
1. STD Direct Flow Chip kit (PCR	24 tests	MAD-003938M-P
STD Flow Chip kit (PCR Mix)	3 strips x 8 tubes (blue)	MAD-003938M-MIX
2. STD Chips	24 tests	MAD-003938M-CH-HS
3. Flow Chip Hybridization Reagents Type I (Manual)	24 tests	MAD-003925M-HS12
Hybridization Solution (Reagent A)	40 ml	MAD-003930MA-HS12-24
Blocking Solution (Reagent B)	10 ml	MAD-003930MB-HS12-24
Streptavidin-Alkaline Phosphatase (Reagent C)	10 ml	MAD-003930MC-HS12-24
Washing Buffer I (Reagent D)	35 ml	MAD-003930MD-HS12-24
Reagent E	10 ml	MAD-003930ME
Washing Buffer II (Reagent F)	18 ml	MAD-003930MF-HS12-24

Table 1. Reagents provided in the STD Direct Flow Chip kit (Manual) format.

KIT/COMPONENTS	FORMAT	REFERENCES
STD Direct Flow Chip kit (Auto)	24 tests	MAD-003938M-HS
1. STD Direct Flow Chip kit (PCR	24 tests	MAD-003938M-P
STD Flow Chip kit (PCR Mix)	3 strips x 8 tubes (blue)	MAD-003938M-MIX
2. STD Chips	24 tests	MAD-003938M-CH-HS
3. Flow Chip Hybridization Reagents Type I (Auto)	24 tests	MAD-003925M-HS
Hybridization Solution (Reagent A)	60 ml	MAD-003930MA-HS24-24
Blocking Solution (Reagent B)	10 ml	MAD-003930MB-HS24-24
Streptavidin-Alkaline Phosphatase (Reagent C)	10 ml	MAD-003930MC-HS24-24
Washing Buffer I (Reagent D)	35 ml	MAD-003930MD-HS24-24
Reagent E	10 ml	MAD-003930ME-HS24

Table 2. Reagents provided in the STD Direct Flow Chip kit (Auto) format.

- Both presentations include 60mL of DNase/RNase-free double distilled water for the handling of clinical samples: RNASE/DNASE-FREE DISTILLED WATER; Ref: MAD-DDW"
- **STD Flow Chip kit (PCR Mix)** contains the monotest lyophilized PCR mix made up of PCR buffer, dNTPs (U/T), DNase/RNase-free water, biotinylated primers, *Hot Start Polymerase* and *Uracil DNA Glycosylase*.





The primers included are specific for the amplification of the following species of pathogenic organisms: *Chlamydia trachomatis, Haemophilus ducreyi,* Herpes simplex virus 1, Herpes simplex virus 2, *Mycoplasma genitalium, Mycoplasma hominis, Neisseria gonorrhoeae, Treponema pallidum, Trichomonas vaginalis* and *Ureaplasma urealyticum/parvum*. Furthermore, the PCR mix includes primers for the amplification of a human genomic DNA fragment used as an internal control, and primers along with DNA of an exogenous control of amplification.

- **STD Chips:** The kit includes a total of 24 Chips or membranes (ref: MAD-003938M-CH-HS) that contain an array of DNA probes specific to each of the pathogens included in the analysis, as well as others corresponding to the internal controls included in this kit. The position of all them on the Chip can be referred to in the section 10 of this manual (INTERPRETATION OF RESULTS).
- Flow Chip Hybridization Reagents: It contains all the reagents necessary for the reverse Flow-Through hybridization process.

# 4. ADDITIONAL REQUIRED MATERIAL NOT SUPPLIED

# 4.1. Reagents and materials

- A. Common reagents to manual and automatic platforms:
  - Disposable gloves.
  - DNasa/RNasa-free tubes of 0.2/0.5 ml
  - Pipette tips with DNasa/RNasa-free filters.
  - For the manipulation of clinical samples of semen: Paraffin Tissue Processing Kit (MAD-003952M).

# B. Specific reagents (Auto, ref: MAD-003938M-HS):

• Washing Reagent (Ref: MAD-003930WSH).

# 4.2. Equipment

# A. Common equipment for the manual and automatic platforms:

- Microcentrifuge.
- Automatic micropipettes: P1000, P200, P20 and P2.
- For enzymatic lysis of semen samples, it is recommended to use a dry-heat block for 1.5mL tubes with a shaking program.
- HybriSoft software.

# B. Specific equipment:

- STD Direct Flow Chip kit (Manual) (Ref: MAD-003938M-HS12):
  - $\circ$  Manual equipment for hybridization hybriSpot 12 (VIT-HS12).
  - o Thermocycler
  - $\circ$   $\;$  Thermal block to heat PCR tubes (can be substituted by a thermocycler)
  - Cold plate (4°C)
  - Thermostatic bath / heater.





- STD Direct Flow Chip kit (Auto) (Ref: MAD-003938M-HS):
  - Automatic equipment for hybridization hybriSpot 24 (VIT-HS24) or hybriSpot 12 PCR AUTO (VIT-HS12a).
  - Thermocycler (not necessary for hybriSpot 12 PCR AUTO).
  - Thermal block to heat PCR tubes (not necessary for hybriSpot 12 PCR AUTO).
  - Cold plate (4°C).

# **5. STORAGE AND STABILITY CONDITIONS**

The *STD Direct Flow Chip* kit consists of three components that are supplied in separate boxes:

- <u>STD Direct Flow Chip kit (PCR Reagents)</u>: Shipment at 2-8 °C\*. Upon receipt, they must be stored at -2-8 °C. They will be stable until the specified expiration date. The PCR reagents must be stored in areas free of DNA or PCR products contamination. Once the package containing the tubes strip with the lyophilized PCR mix is opened, store the remaining tubes up to a maximum of one week at 2-8°C in the original package.
- <u>STD Chips:</u> Sent and stored between 2-8°C\*. <u>Do not freeze</u>. The Chips are stable until their indicated expiration date.
- <u>Hybridization reagents</u>: Sent and stored between 2-8°C\*. <u>Do not freeze</u>. The hybridization reagents are stable until their indicated expiration date. Previous recommendations on the hybridization reagents:
  - The hybridization reagent A must be preheated at 41°C in a thermostatic bath or heater (only for manual format) before use.
  - The rest of the hybridization reagents must be used at room temperature (20-25°C).

Previous chip considerations:

• Once the packaging containing the chips has been opened, keep the cylindrical foam and sorbent packet inside until end of use to ensure the adequate preservation of the membranes.

\*A temperature indicator is included in the package to control the conditions during the shipment. In case the cold chain is interrupted, it is recommended to contact the manufacturer before using the reagents.

# 6. WARNINGS AND PRECAUTIONS

- Read the instructions of use before using this product.
- The safety and disposal precautions are described in the Safety Data Sheet of this product. This product is only intended for professional laboratory purposes, and it is not intended for pharmacological, home or any other type of use. The current version of the Safety Data Sheet of this product can be downloaded in the web page <a href="http://www.vitro.bio">www.vitro.bio</a> or requested at <a href="http://www.vitro.bio">regulatory.md@vitro.bio</a>.
- STD Direct Flow Chip kit uses as a starting material, nucleic acids previously extracted and purified or clinical samples that require a previous manipulation for their analysis. Manipulation protocols are provided in the different types of clinical samples, whose processing has been validated with this kit (see section 7.1).



Vitro S.A. Calle Luis Fuentes Bejarano nº 60. Ed. Nudo Norte Local 3. 41020 Sevilla (Spain). Tel: +34 954 933 200. vitro@vitro.bio; www.vitro.bio





- General considerations to avoid the contamination with PCR product: The greatest contamination source is normally the same amplified PCR product. Therefore, it is recommended to carry out the handling of the amplified products in a different area than the one the PCR reaction is performed. It is recommended to work on different pre- and post-PCR areas where the handling of the test DNA and preparation of the PCR tubes (pre-PCR) and the handling and hybridization of the amplified products (post-PCR) are performed. These areas must be physically separated and different laboratory material must be used (laboratory coats, pipettes, tips, etc.) to avoid the contamination of the samples with the amplified DNA, which could lead to false positive diagnosis. The workflow must always go in a single direction, from the pre-PCR area to the post-PCR area and never the opposite way. The material and personal flow from the post-PCR area to the pre-PCR area must be avoided. Furthermore, in order to avoid the contamination with previous PCR products, the enzyme Uracil-DNA Glycosylase (Cod-UNG), which degrades the PCR products containing dUTP, is included in the kit. It is recommended to include negative amplification controls containing all the reagents handled in the kit, from the extraction to the amplification, except for the DNA sample, in order to detect and control any possible contamination of the reagents with test samples or amplified products. The hybridization in membrane of this control must be negative, marking only the hybridization control and the amplification exogenous control. This way, it is verified that there is no contamination of DNA of patients and/or amplified DNA in the pre-PCR area.
- **Warning**: the use of ethylene oxide for the preparation of clinical samples and/or the PCR mix could interfere in the correct development of the PCR reaction. It is recommended to avoid the use of this component for these purposes.

POTENTIAL WASTES GENERATED AFTER USING THIS PRODUCT	ELW CODE*	TYPE OF WASTE ACCORDING TO ELW*
<ol> <li>Rubbish/Waste generated from hybridization reagents</li> <li>Disposal of Liquid Wastes ("Wastes" in the manual and automatic platforms)</li> </ol>	161001	"Aqueous liquid wastes containing dangerous substances" after adding 10% of the total volume of a disinfectant agent. If the disinfection is not carried out, these wastes must be considered as "wastes whose storage and disposal is subjected to special requirements in order to prevent infection"
<ol> <li>Chips used</li> <li>Perishable material (tubes, tips, aluminum foil, etc.)</li> <li>Any element that has been in contact with DNA</li> </ol>	180103	"Wastes whose collection and disposal is subject to special requirements in order to prevent infection"
<ol> <li>Container for reagents used classified as dangerous (according to the Safety Data Sheet)</li> </ol>	150110	"Containers having residues of or contaminated by dangerous substances"

Table 3. Classification of wastes generated by this kit according to the European Legislation. \*ELW: English acronym for *European Legislation of Waste*.

• **Waste disposal:** The handling of wastes generated by the use of the products commercialized by VITRO, S.A., must be performed according to the applicable law in the country in which these products are being used. As reference, the following table indicates the classification of wastes generated by this





kit according to the European Law, specifically according to the *European Commission Decision of December 18 2014* amending decision 2000/532/CE on the list of waste pursuant to Directive 2008/98/EC of the European Parliament and of the Council:

\*Note: This classification is included as a general guideline of action, being under the final responsibility of the user the accomplishment of all the local, regional and national regulations on the disposal of this type of materials.

# 7. PREPARATION OF THE CLINICAL SAMPLE FOR ANALYSIS

The STD Direct Flow Chip kit has been validated for its use in direct PCR, without previous DNA extraction, from different types of clinical samples. According to the type of starting clinical sample, the processing protocol is the following:

# 7.1. Clinical samples of urine

The *STD Direct Flow Chip kit* has been validated for its use in **direct PCR from clinical urine samples** without being necessary the extraction of DNA. The recommended protocol for the processing of this type of sample is the following:

- Homogenize the urine sample by using a vortex. Take a volume of 400  $\mu$ l of the homogenized mix and transfer it to a suitable microcentrifuge tube.
- Centrifuge the sample for 3 min at 12,000 rpm. After the centrifugation, dispose carefully the supernatant. It is recommended to use a Pasteur pipette or micropipette of 1 mL.
- Wash the cell pellet with 400 μL **DNase/RNase-free double distilled water.** Centrifuge the sample for 3 min at 12,000 rpm. Remove carefully the supernatant.
- Resuspend the cell button in 400 μl **DNase/RNase-free double distilled water** to obtain a homogeneous suspension of cells.
- Shake the sample with a vortex and use 30 ul of the cell suspension as DNA template for the PCR reaction. If the processing analysis is not going to be performed, after that, store at -20°C for a maximum period of 2 months.

# 7.2. Clinical samples of semen

The *STD Direct Flow Chip* kit has been validated for its use in **direct PCR from clinical semen samples** without being necessary the extraction of DNA. The recommended protocol for the processing of this type of sample is the following:

- Homogenize the clinical specimen by using a pipette. Take a volume of 400 µl of the homogenized one and transfer it to a suitable microcentrifuge tube of the same volume.
- Centrifuge the sample for 3 min at 12,000 rpm. After the centrifugation, dispose carefully the supernatant. It is recommended to use a Pasteur pipette or micropipette of 1 mL.
- The clinical samples of semen require an enzymatic treatment for a correct cellular lysis. For this, it is necessary to use the reagents DNA Release and Extraction Buffer included in the Paraffin





*Tissue Processing Kit* (ref: MAD-003952M, property of Vitro, S.A) as follows: add 100  $\mu$ L of lysis solution to each semen sample [dilution 1:50 of *DNA Release* in *Extraction Buffer* (2  $\mu$ L of *DNA Release* solution in 98  $\mu$ L of *Extraction Buffer*)].

- Add 100 µL of lysis solution to the cellular sediment and homogenize.
- Incubate the homogenized product at 55°C for 30 minutes. If possible, incubate the samples with shaking at 1,000 rpm.
- Incubate the cellular lysis at 95°C for 10 mins and after that centrifuge at 2,000 rpm for 1 min to precipitate/sediment the cell debris. Transfer the supernatant to a new tube. Add 27 μl of DNase/RNase-free double distilled water and 3 μl of supernatant as DNA template for the PCR per lyophilized PCR tube. If the processing analysis is not going to be performed, after that, store at 20°C for a maximum period of 2 months.

# 7.3. Urethral, cervical, anal and pharyngeal clinical samples

The *STD Direct Flow Chip* kit has been validated for its use in **direct PCR from clinical urethral, cervical, vaginal, anal and pharyngeal samples** without being necessary the extraction of DNA. All the samples included in this section share the use of pellets to take them. The recommended protocol for the processing of this type of sample is the following:

- Shake the swab in 400 µl DNase/RNase-free double distilled water in a tube of 1.5-2 ml.
- Shake the sample with a vortex and use 30 μl of the cell suspension as DNA template for the PCR reaction.
- If the pellet already contains a transport medium (PBS or the like), it is recommended to follow the protocol below:
  - Shake in vortex to disperse the cells into the solution and transfer the homogenized sample (cell suspension) to a suitable microcentrifuge tube.
  - Centrifuge for 3 min at 12,000 rpm.
  - After the centrifugation, dispose carefully most of the supernatant with a Pasteur pipette or micropipette of 1 mL.
  - Resuspend the cell button in 400 μl DNase/RNase-free double distilled water to obtain a homogeneous cell suspension.
  - $\circ~$  Shake the sample with a vortex and use 30  $\mu l$  of the cell suspension as DNA template for the PCR reaction.
- Note: If the sample is not to be analyzed directly, it must be stored at 4°C for a maximum period of 1 week or -20°C for a maximum period of 2 months.
- Note: For anal clinical samples, an additional wash with DNase/RNase-free double distilled water may be necessary, depending on the turbidity caused by the organic material, in order to minimize the inhibitory effect on the PCR by certain agents contained in this type of samples.

# 7.4. Liquid-based endocervical samples

- Take 400  $\mu$ l of homogenized sample with vortex and put in a tube of 1.5-2 ml.
- Centrifuge for 3 min at 12,000 rpm.





- After the centrifugation, dispose carefully most of the supernatant with a Pasteur pipette or micropipette of 1 mL.
- Resuspend the resulting cell button in 400 µl DNase/RNase-free double distilled water.
- Centrifuge for 3 min at 12,000 rpm. Remove carefully most of the supernatant with a Pasteur pipette or micropipette of 1 mL.
- Resuspend the cell button in 400 μl **DNase/RNase-free double distilled water** to obtain a homogeneous suspension of cells.
- Shake the sample with a vortex and use 30 ul of the cell suspension as DNA template for the PCR reaction.

# 7.5. Considerations for a correct manipulation of clinical samples

Besides carefully follow the indications given by the protocols of processing for clinical specimens (included in this section) a set of incidents that can occur when handle samples and how to avoid them is described below:

INCIDENCES	CONSEQUENCES	HOW TO AVOID IT
Use a different aqueous solution to DNase/RNase-free double distilled water	PCR inhibition	Use DNase/RNase-free double distilled water only
Cells decant on the bottom of the tube.	When pipetting the sample to be added to the PCR, there is a risk of not collecting enough sample.	Shake the tube containing the sample to resuspend the cells before aspirating them with the pipette to add them to the PCR tube. The user must confirm it visually.
The sample contains a lot of cells that form lumps.	When pipetting a lump, the tip can get clogged and no sample can be aspirated to be added to the PCR tube.	The user must confirm visually that the sample is aspirated properly.
The sample contains few cells.	If the cell pellet is resuspended in a large volume of double-distilled water, it may give false-negative results or "Blank" samples due to insufficient material.	Resuspend the cell pellet with less volume of double-distilled water.
The sample was added to the PCR tube, but the tube was not placed in the thermocycler afterwards.	The cells begin to lyse and release proteases that can destroy the polymerase giving "blank" sample results.	Once the samples have been added to the PCR tubes, amplify immediately.
The samples have been stored in double-distilled water for more than 1 week at 4°C.	The cells begin to lyse and release proteases that can inhibit the polymerase giving "blank" sample results.	Use the resuspended samples in double- distilled water after obtaining them. Freeze them if they are not going to be used right after.

Table 4. List of incidents that may occur during the processing of clinical samples, consequences, and solutions offered to obtain an optimal analysis performance.





IMPORTANT: If the clinical samples are not to be processed immediately upon receipt, it is • recommended to store them at 4°C for a maximum period of 1 week or at -20°C for a maximum period of 2 months.

STD Direct Flow Chip kit has also been validated with DNA as the starting material, obtained using the following DNA purification kits and extraction instruments\*:

EXTRACTION KITS	EXTRACTION INSTRUMENTS
MagNA Pure Compact Nucleic Acid Isolation kit I (Roche Diagnostic's)	MagNA Pure Compact Instrument. Version 1.1.2 (Roche Diagnostic's)
EZ1 DSP Virus Kit (Qiagen)	Qiagen EZ1 Advanced XL (Qiagen)

Table 5. Extraction kits and instruments used for the purification of DNA from clinical samples.

\*Note: The system has not been validated with other DNA extraction systems. Therefore, the reliability of the results obtained cannot be guaranteed if a different purification system is used.

# 8. ANALYSIS PROCEDURE FOR HS12 AND HS24 PLATFORMS

# 8.1. Multiplex DNA amplification reaction

The following thermocyclers have been validated with STD Direct Flow Chip:

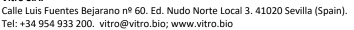
- Veriti 96 Well Thermal Cycler (Life Technologies)
- SimpliAmp Thermal Cycler (Applied Biosystem) -
- TProfessional ThermoCycler (Biometra)

The PCR reaction is carried out in a final volume of 30  $\mu$ l in tubes containing the lyophilized PCR reaction mix.

Procedure:

- Take a tube containing the lyophilized PCR mix per sample to be analyzed. •
- Add 30  $\mu$ l of direct sample in each tube following the recommended protocol in section 7.
- If it involves a semen sample, add 27 μl DNase/RNase-free double distilled water and 3 μl of the • processed clinical sample per reaction tube according to the guidelines given in section 7.
- Homogenize the mix by pipetting and centrifuge for a few seconds.
- If the number of samples to be analyzed is lower or higher than 8, the necessary tubes can be separated from the strip with no need for using complete strips. The rest of the tube strip with the lyophilized PCR mix that is not going to be used at that moment must be stored for maximum of 1 week at 4°C in its original package.
- Place the tubes in the thermocycler and set the following amplification conditions:

PCR PROGRAM				
25°C	10 min	1 cycle		
95°C	3 min	1 cycle		
95°C	30 sec			
55°C	45 sec	40 cycles		







72°C	30 sec		
72°C	5 min	1 cycle	
8°C	∞		
Table 6. PCR program.			

Keep the tubes refrigerated at 8-10 °C when the reaction is finished. If the samples are not going to be processed at that moment, they can be stored in the post-PCR zone at 8-10°C for 1-2 days. To store them for a longer period of time, it is recommended to do so at -20°C.

Important note: If purified DNA is used for PCR, 30  $\mu$ l of this DNA can be added directly to the lyophilized PCR tube.

# 8.2. Flow-through reverse hybridization

All the reagents are provided in a "ready-to-use" format. The membranes are single-use and must be handled with gloves. Depending on the type of kit with which we are working, we will proceed as follows:

# A. For STD Direct Flow Chip kit Manual, (ref: MAD-003938M-HS12):

The full hybridization process is performed semi-automatically in hybriSpot (HS12) following the instructions provided by the wizard of the system. The management of the samples, the capture of images and the analysis and report of the results are performed by the hybriSoft software. Note: Configure the instrument by following the instructions of the user manual (provided with the instrument).

Before starting the hybridization process:

- 1. Denature the PCR products by heating at **95 °C during 10 min** in a thermocycler and **cool quickly in ice** during at least **2 min**.
- 2. Preheat the Reagent A (Reagent A) at 41 °C.
- 3. Place every STD Chip in the position indicated in the platform (HS12).

# Manual hybridization protocol:

- a) Set the temperature of the equipment at 41°C. Add **300**  $\mu$ I of **Reagent A (Hybridization Solution)** preheated for at least 20 minutes at 41°C for every Chip and incubate for at least **2 min at 41°C**.
- b) Remove the **reagent A (Hybridization Solution)** by activating the vacuum pump.
- c) Mix 30 μl of each PCR sample (previously denatured and kept in ice) with 270 μl of Reagent A (Hybridization Solution) (41°C) and dispense the mix on the corresponding STD Chip.
- d) Incubate at **41°C** for **8 min**.
- e) Activate the pump for at least 30 s to remove the PCR products.
- f) Wash **3** times with **300 μl** with **Reagent A (Hybridization Solution)** (41 °C).
- g) Set the temperature at **29°C**.
- h) Add 300  $\mu l$  of Reagent B (Blocking Solution) and incubate for 5 min.
- i) Activate the pump to remove the reagent B.
- j) When the temperature reaches **29°C**, add **300 μL** of **Reagent C (Streptavidin-Alkaline Phosphatase)** to each chip.

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- k) Incubate for 5 min at 29 °C.
- I) Activate the pump to remove the reagent.
- m) Set the temperature at **36°C**.
- n) Wash the membranes **4** times with **300** µl with reagent D (Washing buffer I).
- o) When the temperature has reached 36°C, add 300 μl of Reagent E (developer solution) to every Chip. Incubate for 10 min at 36 °C.
- p) Activate the pump to remove the reagent.
- q) Wash the membranes 2 times with 300 μl with reagent F (Washing buffer II).
- r) Activate the pump to remove the reagent.
- s) Perform the image capture, analysis and result report following the instructions of the equipment's user manual.

# B. For STD Direct Flow Chip kit Auto, (ref: MAD-003938M-HS):

The whole hybridization process is performed automatically on hybriSpot 24 (HS24). The management of the samples, the capture of images and the analysis and report of the results are performed through the hybriSoft software.

Note: Configure the instrument by following the instructions of the user manual (provided with the instrument). Before starting the hybridization process:

- Denature the PCR products by heating at 95 °C during 10 min in a thermocycler and cool quickly in ice during at least 2 min.
- Place the PCR tubes, the STD Chips and the reagents in their corresponding positions of hybriSpot 24.
- 3. Select the corresponding protocol in the equipment to start the automatic process.

# 9. ANALYSIS PROCEDURE FOR HS12a

The amplification through PCR and hybridization processes are performed automatically in the HS12a platform.

The samples processing, images capture and the results analysis are performed by the hybriSoft software.

Before starting the process, it is recommended to carefully read the user manual (included in the HS12a equipment) and follow the instructions to place the strip tubes, chips and hybridization reagents in the instrument.

Procedure:

- Take a tube containing the lyophilized PCR mix per sample to be analyzed.
- Add the DNA samples to a PCR tube following the instructions described in section 8.1.
- Homogenize the mix by pipetting and centrifuge for a few seconds.
- If the number of samples to be analyzed is lower or higher than 8, the necessary tubes can be separated from the strip with no need for using complete strips. The rest of the lyophilized tube strip that is not going to be used at that moment must be stored for maximum of 1 week at 4°C in its original package.





• Follow the instructions in the manual to place the strip tubes, chips and hybridization reagents in the instrument and start the process.

# **10. QUALITY CONTROL PROCEDURE**

The STD Direct Flow Chip kit contains different internal controls to control the quality of the results.

SPOTS	CONTROL	POSITION (see Figure 1)
В	Hybridization control	1A-1B-2I-5E-8A
CI	Exogenous amplification control	1C-5F
BG	Endogenous amplification control	1D-5G
	Table 7. Cantual much as included in CTD C	

Table 7. Control probes included in STD Chip.

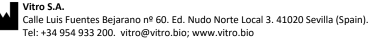
**Hybridization control:** After the development of the membranes, an intense signal must appear in all five hybridization control positions (B), which serve as a quality control. This signal indicates that the hybridization reagents and developing have worked properly. If the signal does not appear, it indicates that an error has occurred during the hybridization process or that a reagent has not been used properly. Furthermore, this signal allows the software to orientate correctly the probe panel to perform the subsequent analysis.

**Exogenous amplification control (CI):** probe for the detection of synthetic DNA included in the PCR mix. This DNA will be co-amplified together with the genetic material of the sample. Two positive signals in the exogenous amplification control (CI) will indicate that the PCR reaction has worked correctly. A negative result in this control does not invalidate the result if the endogenous control has correctly amplified and/or the sample has been positive for any of the organisms included in the panel.

**Endogenous amplification control (BG):** probe for the detection of human beta-globin gene DNA amplified during the PCR. All the samples where the test DNA has been amplified correctly will have a positive signal in the endogenous amplification control (BG). This signal shows the quality/quantity of the DNA used in the amplification. A positive signal indicates that the amplification has worked correctly and that the quality and quantity of the DNA used for it have been optimal. The lack of signal for this control indicates errors during the amplification, due to low quality/quantity of the DNA used in the amplification or lack of human DNA in the amplification. A negative result in this control does not invalidate the result if the exogenous control has correctly amplified and/or the sample has been positive for any of the organisms included in the panel. The latter case is likely to occur with clinical specimen types containing a lower number of human cells.

When a sample is positive for any of the pathogens included in the kit, with a negative result for the exogenous and endogenous amplification controls, the report for the automatic analysis of the results with *HybriSoft software* shows a warning of *"no exogenous control / no human DNA control"* for the user to perform the appropriate verifications before validating the result.

The user is responsible for determining the appropriate quality control procedures for their laboratory and comply with the applicable legislation.







# **11. INTERPRETATION OF RESULTS**

The interpretation of results is done automatically using the analysis software *HybriSoft*. The following scheme shows the arrangement of the probes on the *STD Chip*:

	1	2	3	4	5	6	7	8	9
Α	в	MG				CT-S1		в	
В	В		мн				CT-S2		
с	СІ	τν		UU-P		MG		UU-P	
D	BG		HD				мн		
E		HSV-1/ HSV-2		NG	в	τν		NG	
F			HSV-1		СІ		HD		
G		CTS-1		ТР	BG	HSV-1/ HSV-2		ТР	
н			CT-S2				HSV-1		
I		В							

Figure 1: Scheme of the arrangement of the probes on the array.

"B": Hybridization control

"CI": Exogenous amplification control

"BG": Endogenous amplification control (fragment human ß-Globin)

"X": Specific probes for each pathogen

All the probes are duplicated to guarantee the reliability in the automatic analysis of the results. The hybridization control (B) is repeated in 5 positions and allows the software to orientate correctly the probe panel for its analysis afterwards.

The distribution of the different probes included in the **STD Chip** as well as the possible expected results are shown below:





EXPECTED RESULTS (DETECTED		PROBE/POSITION			
PATHOGENS)	PROBE ID	PROBE	В	CI	BG
Mycoplasma genitalium	MG	2A-6C	1A-1B-2I-5E-8A	/1C-5F	/1D-5G
Mycoplasma hominis	МН	3B-7D	1A-1B-2I-5E-8A	/1C-5F	/1D-5G
Trichomonas vaginalis	τv	2C-6E	1A-1B-2I-5E-8A	/1C-5F	/1D-5G
Ureaplasma urealyticum/parvum	UU-P	4C-8C	1A-1B-2I-5E-8A	/1C-5F	/1D-5G
Haemophilus ducreyi	HD	3D-7F	1A-1B-2I-5E-8A	/1C-5F	/1D-5G
Herpes simplex virus 1*	HSV-1/HSV-2 + HSV-1	2E-6G-3F-7H	1A-1B-2I-5E-8A	/1C-5F	/1D-5G
Herpes simplex virus 2	HSV-1/HSV-2	2E-6G	1A-1B-2I-5E-8A	/1C-5F	/1D-5G
Treponema pallidum	ТР	4G-8G	1A-1B-2I-5E-8A	/1C-5F	/1D-5G
Neisseria gonorrhoeae	NG	4E-8E	1A-1B-2I-5E-8A	/1C-5F	/1D-5G
Chlamydia trachomatis (Serovars A-K)**	CT-S1 + CT-S2	2G-6A-3H-7B	1A-1B-2I-5E-8A	/1C-5F	/1D-5G
Chlamydia trachomatis (Serovars L1-L3)	CT-S2	3H-7B	1A-1B-2I-5E-8A	/1C-5F	/1D-5G
Negative sample			1A-1B-2I-5E-8A	/1C-5F	1D-5G
Negative sample			1A-1B-2I-5E-8A	1C-5F	/1D-5G
Target			1A-1B-2I-5E-8A	1C-5F	
Invalid results			1A-1B-2I-5E-8A		
Hybridization error					

Table 8. Position of the probes on the STD Chip and interpretation of results.

\*Note: The positive identification of the Herpes simplex virus 1 is obtained when, simultaneously, the hybridization signals on the STD Chip from the probe mix HSV-1/HSV-2 (positions 2E-6G) and the probe signal of HSV-1 (positions 3F-7H) occur. When only the hybridization signal of the probe mix HSV-1/HSV-2 (positions 2E-6G) occurs, then it corresponds to the Herpes simplex virus 2. Co-infections of both types of Herpes simplex viruses 1 and 2 show the same patter as an infection by the Herpes simplex virus 1 (2E-6G-3F-7H). Real situations in which only the probe signal HSV-1 on the STD Chip are not contemplated.

\*\*Note: Positive identification of *Chlamydia trachomatis* Serovars A-K is obtained when both probes CT-S1 (2G-6A positions) and CT-S2 (3H-7B) are present in the STD Chip. When only the signal from probe CT-S2 (3H-7B positions) is present, the identification corresponds to Serovars L1-L3 from the same specie. Real situations in which probe CT-S1 is the only one present on the STD chip are not contemplated.

Below, an example of a report in which the analyzed sample has been positive for *Chlamydia trachomatis* (CT) pathogen is shown.





	/itro		STD Direct Flow Chip Kit		it	
	master diagnóstica® Lots					
			PCR:	STD012.2	10/30/2019	
			Chips:	STDE010.6	₽ 9/30/2019	
			Reagent:	HPVH055-2	3/31/2020	
SAMPLE DE	TAILS					
ID SAMPLE:	Sample-36		SA	MPLE TYPE:		
ID PATIENT:		PATIENT:				
SEX:	-	BIRTHDATE:	AG	iE:		
REPORT						

#### STD POSITIVE

Positive sample for:

Chlamydia trachomatis (Serotypes A-K)

The sample is negative for the rest of pathogens included in the STD flow chip test.

#### PROTOCOL

Detection of STD related by PCR and automatic reverse dot blot:

- Herpes simplex virus-1, Herpes simplex virus-2, Neisseria gonorrhoeae, Chlamydia trachomatis (Serovars A-K),

Chlamydia trachomatis (Serovars L1-L3 =Linfogranuloma venereum), Mycoplasma genitalium, Mycoplasma hominis,

Thricomonas vaginalis, Ureaplasmas (urealyticum/parvum), Haemophilus ducreyi and Treponema pallidum.

Clinical Sample processing/DNA purification

- Add processed clinical sample/purified DNA for PCR amplification:

PCR protocol: 1x [25°C, 10 min]; 1x [95°C, 3 min]; 40x [ 95°C, 30 s - 55°C, 45 s - 72°C, 30 s]; 1x [72°C, 5min]; 1x [8°C, ∞].

REVERSE-DOT BLOT protocol:

- Hybridization of the biotinylated PCR products to the STD Direct Flow CHIP.

- Post-hybridization washes.

- Streptavidin-Alkaline Phosphatase incubation.

- NBT-BCIP development.

Automatic analysis of results.

#### NOTES

<b>itro S.A.</b> alle Luis Fuentes Bejara	no nº 60. Ed. Nudo Norte Local 3. 41020 (	Sevilla (Spain).	IVD	Rev.: 2020/01/2
Instr. : Mock	Serial Nº: 100001	hybriSoft:	HSHS 2.2.0.R00 / HSHS IF	PL 1.0.0.R05
Performed by:	Default Tech, tech		Processed:	5/3/2019
FACULTATIVE:	Default Doctor, doctor		Validated:	5/3/2019

Calle Luis Fuentes Bejarano nº 60. Ed. Nudo Norte Local 3. 41020 Sevilla (Spain). Tel: +34 954 933 200. vitro@vitro.bio; www.vitro.bio





Sample-36

\_

PATIENT: BIRTHDATE:

# **STD Direct Flow Chip Kit**

L	0	T:	

PCR:	STD012.2	10/30/2019
Chips:	STDE010.6	₽ 9/30/2019
Reagent:	HPVH055-2	3/31/2020

SAMPLE TYPE:

AGE:

SEX:

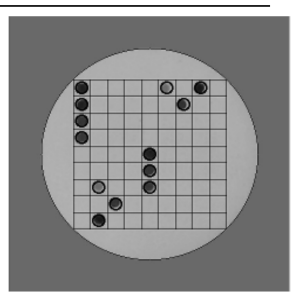
ID SAMPLE:

**ID PATIENT:** 

REPORT

SAMPLE DETAILS

CT-S1 MG в в в МН CT-S2 τν UU-P MG UU-P BG HD мн HSV-1/ NG NG τν HSV-2 HSV-1 HD HSV-1/ CT-S1 ΤР BG ΤР HSV-2 CT-S2 HSV-1 в



- Spot B: Hybridization control (5 signals to orientate the CHIP)

- Spot CI: Amplification control

- Spot BG: DNA Control (Genomic human DNA probe)

- Spot #:Pathogen specific probes

All the spots are printed in duplicate.

#### **ANALYSIS INFORMATION**

Threshold: 4

FACULTATIVE:	Default Doctor, doctor		Validated:	5/3/2019
Performed by:	Default Tech, tech		Processed:	5/3/2019
Instr. : Mock	Serial Nº: 100001	hybriSoft:	HSHS 2.2.0.R00 / HSHS IF	PL 1.0.0.R05

Figure 2. Example of a report of a positive case for Chlamydia trachomatis (CT) pathogen.

#### Vitro S.A.

Calle Luis Fuentes Bejarano nº 60. Ed. Nudo Norte Local 3. 41020 Sevilla (Spain). Tel: +34 954 933 200. vitro@vitro.bio; www.vitro.bio





# **12. PERFORMANCE CHARACTERISTICS**

# **12.1.** Analytical performance on a manual platform (HS12)

# <u>12.1.1. Repeatability</u>

The repeatability of the method was analyzed by using synthetic DNA fragment from each of the specific targets of the pathogens of the panel. Two different concentrations of synthetic DNA were used and from each one of them at least 6 replicates were obtained. The test was performed by the same operator, in a single location and using the same reagents lot.

TARGET	PROBES	No. COPIES / REACTION	POSITIVE / TESTED	% POSITIVE
M. genitalium	MG	50	6/6	100%
M. hominis	MH	10	6/6	100%
U. urealyticum/parvum	UU-P	10	6/6	100%
N. gonorrhoeae	NG	10	6/6	100%
T. vaginalis	TV	10	6/6	100%
HSV-1	HSV-1/HSV-2 + HSV-1	10	7/7	100%
HSV-2	HSV-1/HSV-2	10	7/7	100%
T. pallidum	ТР	10	7/7	100%
H. ducreyi	HD	10	6/6	100%
C. trachomatis (Serovars L1-L3)	CT-S2	10	6/6	100%
C. trachomatis (serovars A-K)	CT-S1 + CT-S2	10	6/6	100%

Table 9. Repeatability assay for each of the pathogens included in the panel.

# 12.1.2. Reproducibility

The method precision was analyzed by simulating the inter-laboratory variability by changing the operator, the PCR mix lot used and the equipment used. 15 clinical cases (10 positive and 5 negative clinical cases) from semen, endocervical and urethral origin were analyzed in parallel. For all the clinical cases, the analysis was performed with the direct method (without previous DNA purification).

	LABORATORY 1				
LABORATORY 2	Positive	Negative	Total		
Positive	15	0	15		
Negative	0	5	5		
Total	15	5	20		

Table 10. Reproducibility test using a clinical sample.

The concordance was calculated for both conditions, obtaining a kappa index of 1.000, a standard error of 0 and a 95% CI of 1.000-1.000. In both conditions, the statistical significance of the reproducibility tests with the *STD Direct Flow Chip* kit were demonstrated.





# 12.1.3. Analytical specificity

Experiments to determine potential cases of cross-reactivity between members of the panel were performed using a specific number of copies of each of the synthetic oligos (1x10<sup>6</sup> copies) representing each pathogen, with no cross-reactivity observed between panel members:

ORGANISM	SPECIFICITY
M. genitalium	100%
M. hominis	100%
U. urealyticum/parvum	100%
N. gonorrhoeae	100%
T. vaginalis	100%
HSV-1	100%
HSV-2	100%
T. pallidum	100%
H. ducreyi	100%
C. trachomatis	100%

Table 11. STD Direct Flow Chip kit intra-panel specificity.

There was also no non-specificity with other pathogens that are either phylogenetically related to the panel members or are associated with other microorganisms that coexist in the same flora:

TESTED MICROORGANISMS (1x10 <sup>4</sup> TOTAL COPIES)				
Aeromonas hydrophila	Listeria monocytogenes			
Acinetobacter baumannii	Morganella morganii			
Campylobacter jejuni	Proteus penneri			
Candida albicans	Providencia rettgeri			
Citrobacter freundii	Providencia stuartii			
Citrobacter koseri	Salmonella enterica			
Citrobacter spp	Salmonella enteritidis			
Cytomegalovirus	Serratia marcescens			
Cryptococcus neoformans	Serratia urealytica			
Enterobacter aerogenes	Staphylococcus aureus			
Enterobacter cloacae	Stenotrophomonas maltophilia			
Enterococcus faecalis	Streptococcus agalactiae			
Epstein Barr virus	Streptococcus pneumoniae			
Escherichia coli	Proteus mirabilis			
Haemophilus influenzae	Pseudomonas aeruginosa			
Klebsiella pneumoniae	Human Papillomavirus (6, 11, 16, 18)			
Kluyvera ascorbata	Yersinia enterocolitica			
Neisseria meningitidis				

Table 12. List of pathogens included in the "inter-panel" Specificity tests by the *STD Direct Flow Chip* kit.





# 12.1.4. Analytical sensitivity

In order to analytically verify the proper performance of the system designed, synthetic double-stranded DNA fragments mimicking the different target regions of the pathogens included in the panel were used. The kit's limit of detection (LoD) was calculated for each one of the analyzed genes. The determination of the minimum number of copies detected was performed through serial dilutions of the synthetic DNA of each one of the pathogens included in the panel with 5 ng of human genomic DNA. In order to calculate sensitivity, each case was repeated 12 times. All the PCRs were hybridized by using the manual platform. The results were analyzed with *HybriSoft software* and the value established for a positive signal was 4 (gray intensity).

ORGANISM	PROBE	No. COPIES/ REACTIO N	POSITIVES/ TESTED	SENSITIVITY	95% CONFIDENCE INTERVAL	SPECIFICITY	95% CONFIDENCE INTERVAL
M. genitalium	MG	50	12/12	100%	69.87-100%	100%	98.10-100%
M. hominis	МН	10	12/12	100%	69.87-100%	100%	98.10-100%
U. urealyticum/ U. parvum	UU-P	10	12/12	100%	69.87-100%	100%	98.10-100%
N. gonorrhoeae	NG	10	12/12	100%	69.87-100%	100%	98.10-100%
T. vaginalis	TV	10	12/12	100%	69.87-100%	100%	98.10-100%
HSV-1	HSV-1/HSV-2 + HSV-1	10	12/12	100%	69.87-100%	100%	98.10-100%
HSV-2	HSV-1/HSV-2	10	12/12	100%	69.87-100%	100%	98.10-100%
T. pallidum	ТР	10	12/12	100%	69.87-100%	100%	98.10-100%
H. ducreyi	HD	10	12/12	100%	69.87-100%	100%	98.10-100%
<i>C. trachomatis</i> (serovars A-K)	CT-S1+CT-S2	10	12/12	100%	69.87-100%	100%	98.10-100%
<i>C. trachomatis</i> (Serovars L1-L3)	CT-S2	10	12/12	100%	69.87-100%	100%	98.10-100%

Table 13. Analytical sensitivity and specificity test using different synthetic DNA copy numbers corresponding to the pathogens included in the panel, establishing a positivity cut-off value of 4.

# 12.2. Analytical performance on the automatic platform HybriSpot 24

The performance and robustness of the *STD Direct Flow Chip* kit in the automatic platform was validated by analyzing a number of limit copies of synthetic DNA fragments from four pathogens included in the panel. The reproducibility of the results obtained by the automatic platform was evaluated by comparing the results obtained in the manual platform. Two types of test were performed:





# 12.2.1. Reproducibility of results in program for a different number of samples

Replicates of a positive sample containing a limit number of *Chlamydia trachomatis* DNA copies (10 copies) were analyzed. These replicates were placed in different positions of the reaction chamber in the equipment and four different protocols were assessed:

- Protocol for 2 samples (2 replicas)
- Protocol for 12 samples (3 replicas)
- Protocol for 15 samples (3 replicas)
- Protocol for 24 samples (4 replicas)

The results were automatically analyzed with *HybriSoft* and differences between the different positions of the reaction chamber nor the used protocol were not detected.

# 12.2.2. Reproducibility of results in different hybridization positions in the automatic platform

Four replicates for three pathogens from the panel were prepared and located in different positions of the two reaction chambers of the equipment, using the protocol for 24 samples. The results were analyzed automatically with *HybriSoft*, showing a percentage of reproducibility for all the analyzed pathogens in different positions.

ORGANISM	No. COPIES/REACTION	POSITIVE/TESTED	DIFFERENCES BETWEEN
Chlamydia trachomatis	10	4/4	No
Trichomonas vaginalis	10	4/4	No
Mycoplasma hominis	10	4/4	No

Table 14. Reproducibility results obtained with the automatic platform and the *STD Direct Flow Chip* kit. The results were analyzed automatically with *HybriSoft* establishing a positivity cut-off value of 4.

This validation proves the reproducibility of the results between the positions 1 and 24 of the equipment and the reproducibility of the results with different programs for a different number of samples.

# 12.3. Analytical performance on the automatic platform HybriSpot 12 PCR AUTO

The performance and the robustness of the STD Direct Flow Chip was validated in the automatic equipment HS12a by analyzing limit concentrations of synthetic DNA fragments of all the pathogens included in the panel. This validation also proves the reproducibility of the results with different programs for different number of samples.

# 12.3.1. Reproducibility of results in program for a different number of samples

Replicates of a positive sample containing a limit number of *Chlamydia trachomatis* DNA copies (10 copies) were analyzed. These replicas were placed in different positions of the reaction chamber of the HS12a system and different protocols were evaluated:

Protocol for 2 samples (2 replicas)





# Protocol for 12 samples (3 replicas)

The results were automatically analyzed with hybriSoft and differences between the different positions of the reaction chamber nor the used protocol weren't detected.

# 12.3.2. Verification of limit of detection

The functioning and the robustness of the STD Direct Flow Chip was validated in the automatic equipment HS12a by analyzing limit concentrations of synthetic DNA fragments of all the pathogens included in the panel.

Three replicas of each positive sample were made. The whole process was performed automatically in two different HS12a equipments, and the results were analyzed with hybriSoft.

Organism	No. copies/ reaction	Positives/ Tested
M. genitalium	50	3/3
M. hominis	10	3/3
U. urealyticum/ U. parvum	10	3/3
N. gonorrhoeae	10	3/3
T. vaginalis	10	3/3
HSV-1	10	3/3
HSV-2	10	3/3
T. pallidum	10	3/3
H. ducreyi	10	3/3
C. trachomatis (Serovars A-K)	10	3/3

Table 15: Verification of sensitivity limit of STD Direct Flow Chip kit in HS12a. The positivity was analyzed with the hybriSoft software by establishing as a cut-off point a value of 4. NT: not tested

# 12.3.3. Clinical performance

**The clinical performance of the STD Direct Flow Chip kit** was validated using DNA purified by means of any of the extraction methods mentioned, as well as using as the starting material for analysis the clinical sample without previous genomic DNA extraction (direct PCR); both are detailed in section 7 of this manual. The diagnostic capacity of the *STD Direct Flow Chip* was evaluated by studying its diagnostic sensitivity and specificity. These two parameters are defined and calculated as follows:

• **Diagnostic specificity** is expressed as a percentage (numerical fraction multiplied by 100), calculated as 100 x the number of true negative values (TN) divided by the sum of true negative values (TN) plus the number of false positive (FP) values, or 100 × TN/ (TN + FP).

• **Diagnostic sensitivity** is expressed as a percentage (numerical fraction multiplied by 100), calculated as 100 × the number of true positive values (TP) divided by the sum of true positive values (TP) plus the number of false negative (FN) values, or 100 × TP/ (TP + FN).





# 12.3.3.1. Diagnostic sensitivity and specificity using purified DNA

Purified DNA from 149 clinical specimens was retrospectively analyzed with *STD Direct Flow Chip* kit. The reference system considered was the standard version, not lyophilized, of the kit.

ORGANISM	TN	FP	тр	FN	DIAGNOSTIC SPECIFICITY	DIAGNOSTIC SENSITIVITY
N. gonorrhoeae	126	0	23	0	100%	100%
T. vaginalis	148	0	1	0	NT	NT
M. genitalium	142	0	7	0	100%	100%
Ureaplasma urealyticum/parvum	88	0	58	3	100%	95.08%
C. trachomatis (serovars A-K)*	78	0	71	0	100%	100%
C. trachomatis (serovars L1-L3)	149	0	0	0	NT	NT
M. hominis	114	0	34	1	100%	97.14%
HSV-1	133	1	15	0	99.26%	100%
HSV-2	145	0	4	0	100%	100%
T. pallidum	148	0	1	0	NT	NT
H. ducreyi**	149	0	0	0	NT	NT

Table 16. Diagnostic sensitivity and specificity results obtained with the *STD Direct Flow Chip* kit from the analysis of purified DNA. NT: Not tested.

\*Note: The 0318 Notified Organism only intervenes in the evaluation of the compliance of the test for *Chlamydia trachomatis* Biovars A-K. The detection of *Chlamydia trachomatis* Biovars L1-L3, as well as the detection of the Biovars A-K in pharyngeal, semen and liquid-based endocervical cytology samples are not regulated by the certification of the 0318 Notified Organism.

\*\*Note: It has not been possible to assess the clinical sensitivity/specificity for *T. vaginalis, T. pallidum* and *H. ducreyi* due to lack of positive clinical samples.

# 12.3.3.2. Diagnostic sensitivity and specificity using clinical samples

The diagnostic specificity and sensitivity of the direct method using clinical samples were determined retrospectively with 141 clinical specimens from different origin, using the standard version of the kit, not lyophilized, as the reference system.

ORGANISM	TN	FP	ТР	FN	DIAGNOSTIC SPECIFICITY	DIAGNOSTIC SENSITIVITY
N. gonorrhoeae	123	0	18	0	100%	94.73%
T. vaginalis	125	0	16	0	100%	100%
M. genitalium	132	1	7	1	99.24%	87.5%
Ureaplasma urealyticum/parvum	96	0	45	0	100%	100%
<i>C. trachomatis</i> (serovars A-K)*	69	0	72	0	100%	100%
C. trachomatis (serovars L1-L3)	140	0	1	0	NT	NT
M. hominis	109	1	31	0	99.09%	100%
HSV-1	140	0	1	0	NT	NT







HSV-2	134	0	7	0	100%	100%
T. pallidum	140	0	1	0	NT	NT
H. ducreyi**	141	0	0	0	NT	NT

Table 17. Diagnostic sensitivity and specificity results obtained with the *STD Direct Flow Chip* kit from the analysis of clinical samples. NT: Not tested.

\*Note: The 0318 Notified Organism only intervenes in the evaluation of the compliance of the test for *Chlamydia trachomatis* Biovars A-K. The detection of *Chlamydia trachomatis* Biovars L1-L3, as well as the detection of the Biovars A-K in pharyngeal samples are not regulated by the certification of the 0318 Notified Organism. \*\*Note: It has not been possible to evaluate the clinical sensitivity/specificity for HSV-1, *T. pallidum* and *H. ducreyi* due to lack of positive clinical samples.

# **13. LIMITATIONS**

Use of inappropriate samples: the method has been validated using clinical specimens directly, or purified genetic material from such samples. The clinical specimen types that have been validated are: urine; sperm; and urethral, endocervical, vaginal, perianal, and pharyngeal swabs. The analysis of any other type of specimen not indicated can lead to wrong or inconclusive results due to PCR reaction inhibition by inhibiting chemical agents.

Problem	Causes	Solutions
	Failure in the hybridization protocol.	
No signal is observed/ there is no hybridization signal		Check that all the reagents have
		been correctly added during the
		hybridization process.
		Check the correct functioning of
		hybriSpot 12/12a/24. Repeat the
		test.
	PCR reagents and/or expired or not	
	stored properly.	Check the expiration date and the
		storage conditions of the reagents
		and the Chips. Repeat the test.
	Chip probes destroyed by rests of	Clean with plenty of distilled water
	decontamination reagents (e.g.	and repeat the experiment.
	Bleach) in the wells.	
	Not enough amount of human DNA	Repeat the PCR by increasing the
No signals in the	in the clinical sample.	amount of starting sample. Repeat
endogenous	in the ennear sample.	the test.
amplification control.		
	Presence of PCR inhibitors.	Purify the DNA of the sample and
		repeat the test.
Presence of	High cell and/or blood content of	Repeat the PCR by diluting the
chromogen	the sample.	starting sample.

# **14. TROUBLESHOOTING**



Vitro S.A.



precipitates in the Chip after finishing		
the hybridization protocol.		
	PCR reagents and/or expired or stored improperly.	Check the expiration date of all the reagents and the storage conditions. Repeat the test.
Weak hybridization signals.	Sample volume used to re-suspend the erroneous lyophilized product.	Repeat the test by using the correct sample volume
	Failure in the hybridization protocol.	Check the correct functioning of hybriSpot HS12/12a/24 and the hybridization protocol. Repeat the test.
	Low quality/quantity of the DNA in the sample.	Concentrate the sample during its processing by adding less water volume.
Absence of exogenous amplification control.	Problems in the amplification by PCR.	Check that the program of the thermocycler is the appropriate, that the mother PCR mix has been prepared properly and that the reagents are stored correctly. Repeat the test.
	Presence of PCR inhibitors in the test sample.	Verify the correct functioning of the extraction system of nucleic acids used. Repeat the test.

Table 18. Possible incidents, causes and solutions against the problems that can arise during the analysis.

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# **16. LABEL SYMBOLS**

Explanation of the symbols of the product label:

$\Sigma$	Expiration date	REF	Catalog number
Ĵ	Temperature limit	LOT	Lot code
***	Manufacturer	ī	Refer to the instructions of use
$\Sigma$	Sufficient content for <n> assays</n>	IVD	Medical product for <i>in vitro</i> diagnosis.

# 17. GLOSSARY

DNA: Deoxyribonucleic acid.
CT: Chlamydia trachomatis.
Cod UNG: Cod Uracil-DNA Glycosylase.
DNase: Deoxyribonuclease.



Calle Luis Fuentes Bejarano nº 60. Ed. Nudo Norte Local 3. 41020 Sevilla (Spain). Tel: +34 954 933 200. vitro@vitro.bio; www.vitro.bio





dUTP: Deoxyuridine Triphosphate. **STD:** Sexually transmitted disease. FN: False negative. False negative results. FP: False positive. False positive results. HD: Haemophilus ducreyi. HS12: HybriSpot 12 equipment. HS12a: HybriSpot 12 PCR AUTO equipment. HS24: HybriSpot 24 equipment. L1-L3: Chlamydia trachomatis serovars L1-L3. MG: Mycoplasma genitalium. MH: Mycoplasma hominis. NBT-BCIP: Nitroblue Tetrazolium Chloride- 5-Bromo-4-Chloro-3-Indolyl phosphate. NG: Neisseria gonorrhoeae. PCR: Polymerase Chain Reaction. RNase: Ribonuclease. TN: True negative. True negative results. **TP:** *Treponema pallidum.* TP: True positive. True positive results. TV: Trichomonas vaginalis. UP: Ureaplasma parvum. UU: Ureaplasma urealyticum. HSV-1: Herpes simplex virus 1. HSV-2: Herpes simplex virus 2.

