For professional in vitro diagnostic use only.

INTENDED USE

Vitassay Clostridium difficile GDH+Toxin A+B is a rapid, immunochromatographic assay for the simultaneous qualitative detection of *Clostridium difficile* glutamate dehydrogenase (GDH), Toxin A and Toxin B in human stool samples.

Simple, non-invasive and highly sensitivity immunoassay to make a presumptive diagnosis of *Clostridium difficile* infection.

INTRODUCTION

Clostridium (Clostridioides) difficile, is a gram-positive anaerobic enterotoxigenic bacillus identified as one of the leading causes of infectious diarrhoea related to antibiotic treatment (10-15%). It is a major public health concern, as it responsible of broad outbreaks in hospitals and nursing homes. C. difficile is also associated with high mortality rates in the elderly (Marra et al., 2020). The main risk factors for C. difficile infections are age (>65 years-old), hospitalisation, being immunocompromised and/or previous infection (Abt et al., 2016).

C. difficile can be divided into two major strains: nontoxigenic and toxigenic, but only the later one causes disease in humans. These toxigenic strains produce toxin B (TcdB) alone, or in combination with toxin A (TcdA) and the pathogenicity depends on the expression of these two toxins (Burke et al., 2014). The toxins are produced and secreted in the large intestine under limit nutrient availability and undergo endocytosis by the intestinal epithelial cells. The activation of these toxins in the cytosol, produces cell necrosis and the subsequent loss of intestinal membrane integrity. This process leads to host exposure to intestinal microorganisms and the subsequent activation of the inflammatory response (CDC).

Different toxigenic strains have been reported to cause important outbreaks around the globe. In 2003, the new strain NAP1/ribotype 027 was a source of C. difficile epidemics in the US and Canada. It has also been described across Europe and Chile, and t is associated with higher morbidity and recurrent rates. It is of a particular interest since it has a mutation in an inhibitory gene, leading to an increase toxin A and B production (McDonald et al., 2005). In other parts of the world, as Asia, it is rare to find 027 ribotype, however 017 or 002 accounts for a large number of cases (Burke et al., 2014).

Treatment is based on antibiotic administration, including and vancomycin and fidaxomicin and metronidazole/vancomycin. However, antibiotics are not always affective and there is a 25% risk of having recurrent episodes. Innovative treatments, like FMT (faecal microbiota transplant), antibody-based therapy and

microbiota-based drugs have risen. There are two vaccines under evaluation (Guery et al., 2019).

Signs of infection might appear after taking antibiotics or being hospitalised/during hospitalisation. Common symptoms are severe diarrhoea, fever, stomach tenderness, loss of appetite and nausea (CDC). Measurements to avoid spreading must be taking, such as good hygiene and avoiding sharing towels/linen.

Systematic testing for *C. difficile* is recommended in case a diarrhoea happens in a healthcare environment, or where the common causatives of intestinal infections have been discarded.

PRINCIPLE

Vitassay Clostridium difficile GDH+Toxin A+B is a qualitative immunochromatographic assay to make a presumptive diagnosis of Clostridium difficile infection.

Strip A: The test line zone of the nitrocellulose membrane is precoated with monoclonal antibodies against GDH.

Strip B: The test line zone of the nitrocellulose membrane is precoated with monoclonal antibodies against Toxin A of *Clostridium difficile*.

Strip C: The test line zone of the nitrocellulose membrane is precoated with monoclonal antibodies against Toxin B of *Clostridium difficile*.

During the process, the sample reacts with the antibodies against GDH (strip A), Toxin A (strip B) and/or Toxin B (strip C), forming conjugates. The mixture moves upward on the membrane by capillary action. If the sample is positive, the antigens of the diluted sample react with the conjugate complex and a red line will be visible. Although the sample is positive or negative, the mixture continues to move across the membranes and the control line always appears.

The presence of a green line (in the control zone (C)) indicates that sufficient volume is added; proper flow is obtained and serves as an internal control for the reagents.

PRECAUTIONS

- For professional in vitro use only.
- Do not use after expiry date.
- Read the instructions for use carefully before using the test.
- Do not use the kit if the label sealing the outer carton is torn or if the bags are open or damaged on arrival.
- Do not use the tests if the desiccant material is missing or broken inside the aluminium pouch.
- Specimens should be considered potentially hazardous and should be handled in the same manner as an infectious agent, following local/national regulations. A new test should be used for each sample to avoid contamination errors.

VITASSAY

Clostridium difficile GDH+Toxin A+B

Rapid test for the qualitative detection of Clostridium difficile Glutamate Dehydrogenase (GDH), Toxin A and Toxin B in human stool samples.

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- Material exposed to the samples should also be considered potentially hazardous and should be handled in the same manner as an infectious agent, following local/national regulations.
- Do not reuse. This is a single-use device.
- Used material should be disposed of in an appropriate biohazard container after testing.
- The reagents contain a preservative called sodium azide (<0.1%). Avoid any contact with skin or mucous membranes.
- In accordance with current regulations Vitassay Clostridium difficile GHD+Toxin A+B does not contain substances and/or mixtures that are hazardous or present in a concentration above the limits for their declaration. The safety data sheet is available on request (not included).
- All reagents included in the kit are approved for use with Vitassay Clostridium difficile GDH+Toxin A+B only. Do not mix or use the components with other batches of Vitassay. Do not use with reagents from other kits or commercial assays.
- Follow Good Laboratory Practices. These practices should include, but are not limited to, personal protective equipment (PPE), such as lab coat, surgical or appropriate mask or face shield, disposable gloves and eye protection. Take the necessary precautions during sample collection, transport, storage, handling and disposal. Each sample must be correctly and unequivocally identified to ensure proper traceability of samples.
- In case of spillage, clean thoroughly with a suitable disinfectant.
- Do not eat, drink or smoke in the workplace.
- The presence of yellow lines in the result window (control line area and test line area), before using the test, is completely normal and does not imply a failure in the functionality of the test.
- The visual interpretation of the results is done by coloured lines, the interpretation of the results should be done by a professional user without problems of visualisation and colour interpretation.
- A certificate of analysis can be provided on request (not included).

STORAGE AND STABILITY

The storage temperature of the kits should be 2-30°C.

Do not freeze.

Under these conditions, they can be used until the expiry date indicated on the kit label.

All kit components are for single use only and must remain in their primary packaging until use. The test must remain in the sealed pouch until use.

MATERIALS					
MATERIAL PROVIDED	MATERIAL REQUIRED BUT NOT PROVIDED				
10X Vitassay Clostridium difficile GDH+Toxin A+B	 Specimen collection container. Disposable gloves. 				
Instructions for use. 10 X vials with diluent for sample dilution.	Timer. Spatula.				

SPECIMEN COLLECTION

Collect sufficient quantity of faeces: 1-2g or mL for liquid samples. Stool specimens should be collected in clean and dry containers.

If processed the day after collection, samples can be stored at room temperature. However, we recommend testing immediately after sample collection.

Samples can be stored for up to 7 days at 2-8°C, and also frozen at -20°C for up to one year. Samples shall be brought to room temperature before testing.

Ensure that only the necessary quantity is thawed, as freezing freeze-thaw cycles are not recommended. Homogenise stool samples as thoroughly as possible prior to preparation.

SPECIMEN PREPARATION

- 1 Remove the cap of the vial with diluent for sample dilution (figure 1) and use the spoon to collect sufficient sample quantity. For solid stool, insert the spoon in 4 different areas of the stool sample (figure 2), remove any excess sample with a spatula (figure 3), and place the spoon cap back into the vial for sample dilution (figure 4). For liquid stool, take a spoonful of the sample (figure 3) and transfer it into the vial for sample dilution.
- 2.Close the vial for sample dilution tightly and shake it to dilute and mix the sample with the diluent (figure 4).





Ctra. N.330, Km.566 22197-Cuarte (Huesca, SPAIN) www.vitassay.com Vial for sample dilution



Insert the spoon in 4 different areas of the stool.



Remove excess sample with a spatula. Liquid samples: full spoon.





Put the sample into the vial, close the cap and shake.

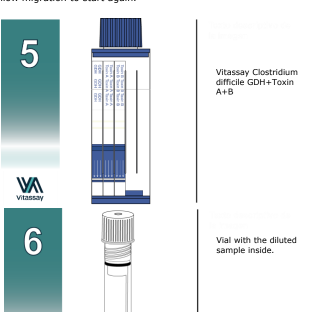


PROCEDURE

Allow the test, stool sample, controls and diluent to reach room temperature (15-30°C) prior to testing. Do not open pouches until the performance of the assay.

- 1. Shake the vial with the sample vigorously to obtain a good sample dilution.
- 2. Remove the **Vitassay Clostridium difficile GDH+Toxin A+B** from its sealed bag just before using it (figure 5).
- 3. Take the vial for sample dilution containing the diluted sample (figure 6), place it inside the multiplex tube (figure 7). Screw the cap of the multiplex tube tightly (figure 8). The bottom of the vial for sample dilution will break and the diluent+sample solution reaches the sample zone of the strips (figure 9).
- 4. Leave the multiplex tube vertically on a flat surface and read the results at **10 minutes**. Do not read the test results later than 10 minutes.

If the test does not run due to solid particles (the sample is not homogenized), migration process can stop on one or more strips. In this case, tap the end of the multiplex tube on hard surface to allow migration to start again.

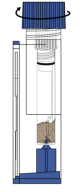






Introduce the vial with the diluted sample into the multiplex.





Close the cap and the bottom of the diluent vial will break.



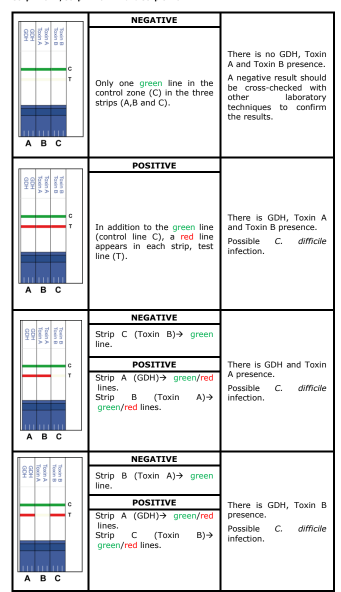
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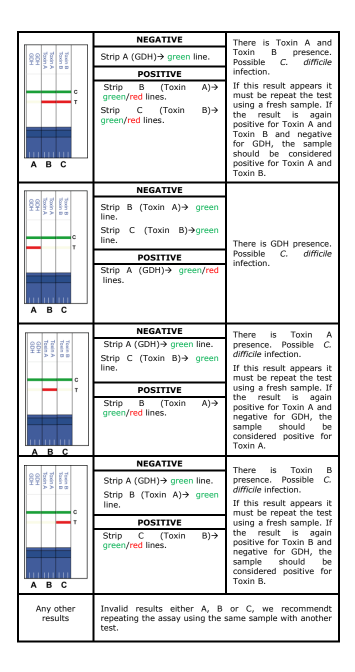


Ctra. N.330, Km.566 22197-Cuarte (Huesca, SPAIN) www.vitassay.com Reaction takes place. Read results at 10 minutes.

INTERPRETATION OF THE RESULTS

Strip A: GDH, Strip B: Toxin A and Strip C: Toxin B





A very low percentage of specimens might result negative for GDH but positive for toxins.

Notes: The intensity of the red coloured test line in the result line region (T) will vary depending on the concentration of antigens in the specimen.

Positive results detailed in the above table should be followed up with additional confirmatory diagnostic procedures.

Invalid results: Total absence of any control coloured lines (green) indicates an invalid result, regardless of the appearance or not of the test lines (red). Wrong procedural techniques or deterioration of the reagents are mostly the main reasons for control line failure. Review the procedure and repeat the assay with a new test. If the problem persists, discontinue using the kit and contact your local distributor.

If the results are read later than 10 minutes after sample addition, they shall be considered invalid.

OUALITY CONTROL

Internal procedural controls are included in **Vitassay GDH+Toxin A+B**. Green lines appearing in the results window are internal controls, which confirm sufficient specimen volume and correct procedural technique.

LIMITATIONS

- Vitassay Clostridium difficile GDH+Toxin A+B test must be carried out within 2 hours of opening the sealed bag.
- An excess of stool sample could cause wrong results (brown bands appear). Dilute the sample with the diluent and repeat the test.
- The intensity of test line may vary depending on the concentration of antigens.
- The use of other samples different from human fecal samples has not been established.
- The quality of Vitassay Clostridium difficile GDH+Toxin A+B depends on the quality of the sample; Proper fecal specimens must be obtained.
- Positive results determine the presence of GDH, Toxin A and/or Toxin B of Clostridium difficile in fecal samples. A positive result should be followed up with additional laboratory techniques (toxigenic culture) to determine the strain. A confirmed infection should only be made by a physician after all clinical and laboratory findings have been evaluated and must be based in the correlation of the results with further clinical observations.
- Negative results should not be considered as conclusive; it is
 possible that the concentration of antigen is lower than the
 detection limit value. If symptoms or situation still persist a
 Clostridium difficile determination should be carried out on a
 sample from an enrichment culture.
- **Bloody stool samples** and/or **mucinous** stool samples can cause non-specific reactions in the test. Such positive samples

- should be followed up with other diagnostic techniques to confirm the result.
- This test is a screening tool. The final diagnosis must be confirmed by a physician after a thorough evaluation of the clinical history and additional confirmatory tests.

EXPECTED VALUES

Clostridium difficile is associated with 95-100% of cases of pseudomembranous colitis, 60-75% of cases of antibiotic-associated colitis and 35% of cases of antibiotic-associated diarrhoea cases. The other causes of antibiotic-associated diarrhoea are largely unknown.

According to ECDC, in 2016-2017, 72.0% of the CDI (C, difficile infection) cases with case-based data were above 64 years old and the majority (56.4%) were female. More than half (n=3 446/5 863: 58.8%) of the CDI cases had had contact with healthcare in the three months before the current healthcare admission, of which the vast majority (n=2 804/3 446; 81.4%) had had contact with a hospital. Fewer were reported to have had contact with a long-term care facility (LTCF) (n=431; 12.5%). There were an estimated 189 526 healthcare-associated (HA) CDI cases (cumulative 95% confidence interval (95%CI): 105 154-340 978) in EU/EEA countries/administrations annually in 2016-2017. In 2016-2017, the crude incidence density of CDI was 3.48 cases per 10 000 patient-days. This was higher in tertiary care hospitals (3.87 cases per 10 000 patient-days) than in secondary or primary care hospitals (3.46 and 2.28 cases per 10 000 patient-days, respectively).

PERFORMANCE CHARACTERISTICS

Clinical Sensitivity and Specificity

Two retrospective evaluations were carried out with stool samples, comparing the results obtained by **Vitassay Clostridium difficile GDH+Toxin A+B** against results obtained with commercial assays based on the immunochromatographic technique, considered as reference methods.

In an initial study, a total of 126 samples from patients with *C. difficile* infection were compared. The results are shown below:

GDH

		Reference method (GDH)		
		Positive	Negative	Total
Vitassay Clostridium	Positive	51	0	51
difficile GDH+Toxin A+B	Negative	0	75	75
GDH	Total	51	75	126

Table 1. Results of Vitassay Clostridium difficile GDH+ Toxin A + B (GDH) compared to a commercial CE-IVD kit.



Vitassay Clostridium difficile GDH + Toxin A + B (GDH) Reference method (GDH)				
	Value	IC 95%		
Sensibility	100%	93.0-100%		
Specificity	100%	95.2-100%		
PPV	100%	93.0-100%		
NVP	100%	95.2-100%		

Table 2. Sensitivity, specificity, positive predictive values and negative predictive values of the Vitassay Clostridium difficile GDH +Toxin A + B (GDH) kit compared to a commercial CE-IVD kit.

Toxin A

		Reference method (Toxin A+B)		
		Positive	Negative	Total
Vitassay Clostridium	Positive	28	0	28
difficile GDH+Toxin A+B	Negative	1	97	98
Toxin A	Total	29	97	126

Table 3. Results of Vitassay Clostridium difficile GDH + Toxin A + B (Toxin A) compared to a commercial CE-IVD kit.

Vitassay Clostridium difficile GDH + Toxin A + B (Toxin A) vs Reference method (Toxin A+B)					
Value IC 95%					
Sensitivity	96.6%	92.2-99.9%			
Specificity	100%	96.2-100%			
PPV	100%	87.7-100%			
NVP	99.0%	94.4-100%			

Table 4. Sensitivity, specificity, positive predictive values and negative predictive values of the Vitassay Clostridium difficile GDH +Toxin A + B (Toxin A) kit compared to a commercial CE-IVD kit.

Toxin B

		Reference method (Toxin A+B)		
		Positive	Negative	Total
Vitassay Clostridium	Positive	36	1	37
difficile GDH+Toxin A+B	Negative	0	89	89
Toxin B	Total	36	90	126

Table 5. Results of Vitassay Clostridium difficile GDH + Toxin A + B (Toxin B) compared to a commercial CE-IVD kit.

Vitassay Clostridium difficile GDH + Toxin A + B (Toxin B) vs Reference method (Toxin A+B)

	95% IC	IC 95%	
Sensitivity	100%	90.3-100%	
Specificity	98.9%	94.0-100%	
PPV	97.3%	85.8-99.9%	
NVP	100%	95.9-100%	

Table 6. Sensitivity, specificity, positive predictive values and negative predictive values of the Vitassay Clostridium difficile GDH +Toxin A + B (Toxin B) kit compared to a commercial CE-IVD kit.

In a second, more recent evaluation, 250 frozen clinical specimen (stool) samples from patients with suspected *C. difficile* infection were evaluated. This test evaluated the performance of the Vitassay kit against another commercial CE-IVD kit and discrepancies were resolved using a certified nucleic acid amplification method.

GDH

		Reference method (GDH)		
		Positive	Negative	Total
Vitassay	Positive	86	1	87
Clostridium difficile	Negative	3	160	163
GDH+Toxin A+B	Total	89	161	250

Table 7. Results of Vitassay Clostridium difficile GDH + Toxin A + B (GDH) compared to a commercial CE-IVD kit.

Vitassay Clostridium difficile GDH + Toxin A + B (GDH) Reference method (GDH)				
	Value	IC 95%		
Sensitivity	96.6%	90.5-99.3%		
Specificity	99.4%	96.6-100%		
PPV	98.9%	93.0-100%		
NVP	98.2%	95.2-100%		

Table 8. Sensitivity, specificity, positive predictive values and negative predictive values of the Vitassay Clostridium difficile GDH +Toxin A + B (GDH) kit compared to a commercial CE-IVD kit.

Toxin A

		Reference method (Toxin A+B)		
		Positive	Negative	Total
Vitassay Clostridium	Positive	51	0	51
difficile Toxin A+B	Negative	1	198	199
Toxin A	Total	52	198	250

Table 9. Results of Vitassay Clostridium difficile GDH + Toxin A + B (Toxin A) compared to a commercial CE-IVD kit.

Vitassay Clostridium difficile Toxin A + B (Toxin A) vs Reference method (Toxin A+B)

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	Value	IC 95%	
Sensitivity	98.1%	89.7-100%	
Specificity	100%	98.2-100%	
PPV	100%	93.0-100%	
NVP	99.5%	97.2-100%	

Table 10. Sensitivity, specificity, positive predictive values and negative predictive values of the Vitassay Clostridium difficile GDH +Toxin A + B (Toxin A) kit compared to a commercial CE-IVD kit

Toxin B

		Reference (Toxin A+B)		
		Positive	Negative	Total
Vitassay Clostridium	Positive	40	0	40
difficile Toxin A+B	Negative	4	206	210
Toxin B	Total	44	206	250

Table 11. Results of Vitassay Clostridium difficile GDH + Toxin A + B (Toxin B) compared to a commercial CE-IVD kit.

Vitassay Clostridium difficile Toxin A + B (Toxin B)vs Reference method (Toxin A+B)			
	Value	IC 95%	
Sensitivity	90.9%	78.3-97.5%	
Specificity	100%	98.2-100%	
PPV	100%	91.2-100%	
NVP	98.1%	95.2-99.9%	

Table 12. Sensitivity, specificity, positive predictive values and negative predictive values of the Vitassay Clostridium difficile GDH +Toxin A + B (Toxin B) kit compared to a commercial CE-IVD kit.

In brief, **Vitassay Clostridium difficile GDH + Toxin A + B** has been tested on a total of 376 clinical faecal samples. The **pooled** analysis of both studies shows an overall clinical **sensitivity** for GDH of 0.97 (0.93-0.99), toxin A of 0.99 (0.97-0.99) and for toxin B of 0.95 (0.87-0.98). On the other hand, the overall clinical **specificity** calculated for GDH is 0.99 (0.97-1), toxin A is 1 (0.98-1) and for toxin B 0.99 (0.98-1).

Altogether, the results show that **Vitassay Clostridium difficile GDH+Toxin A + B** has a high sensitivity and specificity for detecting the enzyme glutamate dehydrogenase (GDH), toxins A and B of *Clostridium difficil*.

Analytical sensitivity

The Vitassay **Clostridium difficile GDH +Toxin A+B** limit of detection (LoD) is 0.39 ng/mL for *C. difficile* Glutamate dehydrogenase (GDH) antigen, 2 ng/mL for *Clostridium difficile* Toxin A and 3.12 ng/mL for Toxin B. Note that a general, reference standard has not been approved; LoD was determined using recombinant GDH (glutamate dehydrogenase) protein, and purified *C. difficile* Toxins A and B as internal standards.



Cross reactivity

No cross reactivity was detected against other gastrointestinal pathogens, other organism, substances and/or faecal markers that are occasionally present in faeces.

GDH

Adenovirus	Escherichia coli 0111	Peptostreptococcus anaerobius
Astrovirus	Escherichia coli 026	Rotavirus
Bovine Haemoglobin	Escherichia coli 0157	RSV
Campylobacter coli	Clostridium difficile Toxin A	Salmonella paratyphi A
Campylobacter jejuni	Giardia	Salmonella typhi
Clostridium bifermentas	Helicobacter pylori	Salmonella typhimurium
Clostridium butyricum	Human Calprotectin	Shigella boydii
Clostridium haemolyticum	Human Haemoglobin	Shigella dysenteriae
Clostridium novyi	Human Lactoferrin	Shigella flexneri
Clostridium tetani	Human Transferrin	Shigella sonnei
Clostridium difficile Toxin B	Influenza A virus	Staphylococcus aureus
Clostridium perfringens	Influenza B virus	Streptococcus pneumococcal
Clostridium septicum	Salmonella enteritidis	Streptococcus pyogenes
Cryptosporidium parvum	Norovirus GII	Yersinia Enterocolitica 0:3
Entamoeba histolytica	Porcine haemoglobin	Yersinia Enterocolitica O:9

It has been observed that *C. sporogenes* (CECT 485) y *C. bolutinium* (CECT 551) can be detected with **Vitassay Clostridium difficile GDH+Toxin A + B.** Intercomparison studies have shown that *C. sordelii* (ATCC 9714) yielded a positive signal in this Vitassay kit, however the origin of this positive result could not be stablished. Hence, it was not considered a genuine cross-reactivity.

Toxin A+B

Adenovirus	Giardia	Salmonella enteritidis
Astrovirus	Helicobacter pylori	Streptococcus pyogenes
Human calprotectin	Bovine haemoglobin	Staphylococcus aureus
Campylobacter coli	Porcine haemoglobin	Human transferrin
Campylobacter jejuni	Human haemoglobin	Salmonella paratyphi A
Clostridium difficile antigen GDH	Influenza A virus	Salmonella typhi
Clostridium difficile Toxin A (strip: Toxin B)	Influenza B virus	Salmonella typhimurium
Clostridium difficile Toxin B (strip: Toxin A)	Human lactoferrin	Shigella boydii
Clostridium perfringens	Legionella pneumophila	Shigella dysenteriae
Coronavirus	Listeria monocytogenes	Shigella flexneri
Cryptosporidium parvum	Norovirus GI	Shigella sonnei

	Entamoeba histolytica	Norovirus GII	Streptococcus pneumococcal
	Escherichia coli 0111	Rotavirus	Yersinia enterocolitica O:3
	Escherichia coli 026	RSV	Yersinia enterocolitica O:9
ı	Escherichia coli O157		

Interferences

Possible interferences were assessed using the following **exogenous compounds and drugs** at the given concentrations (no interferences were found):

Aceltylcysteine (Fluimucil)	3.0 mg/mL	No interference
Acetyl Salicylic (<i>Adiro</i>)	0.3 mg/mL	No interference
Afrin (Oxymetazoline)	5.0·10-2 mg/mL	No interference
Almagato (Almax)	3.0 mg/mL	No interference
Amantadine	0.3 mg/mL	No interference
Ambroxol hydrochloride (Mucosan)	0.3 mg/mL	No interference
Amoxicillin	3.0 mg/mL	No interference
Ampicillin	3.0 mg/mL	No interference
Benzocaine (Angileptol)	0.3 mg/mL	No interference
Biotine	100.0 μg/mL	No interference
Carbocisteine (Iniston mucolítico)	3.0 mg/mL	No interference
Ciprofloxacin	0.3 mg/mL	No interference
Cloperastine (Flutox)	0.3 mg/mL	No interference
Codeine (Toseina)	0.2 mg/mL	No interference
CVS Nasal Drops (Phenylephrine)	10.0 mg/mL	No interference
CVS Nasal Spray (Cromolyn)	4 mg/mL	No interference
Dexchloropheniramine (Polaramine)	0.3 mg/mL	No interference
Dexketoprofen trometamol (Enantyum)	0.3 mg/mL	No interference
Ebastine (Ebastel)	3.0 mg/mL	No interference
Fluticasone Propionate	5.0-10-2 mg/mL	No interference
Fosfamycin (Monurol)	3.0-10-3 mg/mL	No interference
Heparin (Hibor)	350.0 IU/mL	No interference
Hydroxyzine dihydrochloride	0.3 mg/mL	No interference
Homeopathy	1/10 dilution	No interference
lbuprofen (<i>Espidifen</i>)	3.0 mg/mL	No interference
Levofloxacin	3.0 mg/mL	No interference
Loperamide hydrochloride (Fortasec)	0.15 mg/mL	No interference
Loratadine	0.3 mg/mL	No interference
Lorazepam	3.0·10-3 mg/mL	No interference
Lysine Carbocysteinate (Pectox)	3.0·10-2 mg/mL	No interference
Macrogol 3350 (Movicol)	3.0 mg/mL	No interference
Mercaptopurine	0.3 mg/mL	No interference
Metamizole (<i>Nolotil</i>)	5.0 mg/mL	No interference
Metronidazole	3.0 mg/mL	No interference
Mupirocin	2.5·10-2 mg/mL	No interference
Naso GEL	0.9 mg/mL	No interference
Omeprazole	2.0·10-3 mg/mL	No interference

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Oseltamivir	0.3 mg/mL	No interference
Paracetamol (Dolocatil)	5.0 mg/mL	No interference
Phenoxymethylpenicillin potassium	3.0 mg/mL	No interference
Phenulpropanolamine	0.3 mg/mL	No interference
Prednisone	0.3 mg/mL	No interference
Ribavirin	3.0 mg/mL	No interference
Rifampicin (Rifaldin)	0.3 mg/mL	No interference
Sore Throat Phenol spray	0.5 mg/mL	No interference
Tobramycin	0.3 mg/mL	No interference
ZICAM	1.0 mg/mL	No interference

In addition, the following human **endogenous substances** were also analysed. As before, no interferences were found:

Human blood	5.0 %	No interference
Human calprotectin	5.0 μg//mL	No interference
Human haemoglobin	5.0 μg//mL	No interference
Human lactoferrin	5.0 μg//mL	No interference
Human transferrin	0.5 μg//mL	No interference
Mucine	0.5 %	No interference

Repeatability and Reproducibility

Repeatability and reproducibility studies performed with different positive and negative samples yielded no differences between the evaluations.

Hook effect

No inhibitory hook effect was demonstrated at concentrations higher than 10^5 times the limit of detection (10^5 xLoD) for GDH, nor 10^2 times the limit of detection (10^2 xLoD) for toxin A, nor at concentrations 10^4 times the limit of detection (10^4 xLoD) for toxin B in the **Vitassay Clostridium difficile GDH+Toxin A + B** kit.

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SYMBOLS FOR IVD COMPONENTS AND REAGENTS

F09-06 Rev01

IVD	i <i>n vitro</i> diagnostic device	*	Keep dry
(li	Consult instructions for use	1	Temperature limitation
\square	Use by	ш	Manufacturer
LOT	Batch code	\sum_{n}	Contains sufficient for <n> test</n>
DIL	Sample diluent	REF	Catalogue number
CE	CE Marking		





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