

VITASSAY

Norovirus GI+GII

Rapid test for the simultaneous qualitative detection of genogroups I and II of Norovirus (GI and GII) in human stool samples.

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For professional *in vitro* diagnostic use only.

INTENDED USE

Vitassay Norovirus GI+GII is a rapid, immunochromatographic, one step assay for the simultaneous qualitative detection of Norovirus GI and Norovirus GII in human stool samples.

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

INTRODUCTION

Norovirus is a genus within the Caliciviridae family, classified into at least 6 genogroups (GI-GVI) and further into 29 genotypes based on the major viral capsid protein (VP1), including the 3 genogroups that infect humans (GI, GII, and GIV).

In spite of this massive genetic heterogeneity, the vast majority of human Norovirus infections are associated with GII viruses.

Norovirus is a clinically important RNA virus that is estimated to cause almost half of all cases of acute gastroenteritis, globally. Norovirus is also the only human enteric virus known to cause pandemics of acute gastroenteritis. Norovirus infects people of all ages; however, the majority of Norovirus-associated gastroenteritis outbreaks occur within institutional settings, such as aged-care facilities, hospitals and child care centers, thereby affecting the most vulnerable in the community including the elderly, immunocompromised and young children.

Noroviruses can infect humans via multiple routes, including the oral route, transmitted through contact with fecal matter or aerosolized vomitus from infected people, as well as contaminated surfaces, food, or water.

After an incubation period of 12 to 48 h, norovirus illness is characterized by projectile vomiting, nonbloody diarrhea, nausea, abdominal cramps, and low-grade fever. Some persons might experience only vomiting or diarrhea. In healthy individuals, the duration of symptoms is usually not longer than 48 h, and the disease is self-limiting in most patients. However, young children and the elderly are at increased risk for more-severe and prolonged illness leading to hospitalization, while the disease is increasingly recognized as an important cause or chronic gastroenteritis in immunocompromised patients.

PRINCIPLE

Vitassay Norovirus GI+GII is a qualitative immunochromatographic assay for the detection of genogroups I (GI) and II (GII) of Norovirus in human stool samples.

Strip A: The test line zone of the nitrocellulose membrane is pre-coated with monoclonal antibodies against Norovirus GI.

Strip B: The test line zone of the nitrocellulose membrane is pre-coated with monoclonal antibodies against Norovirus GII.

During the process, the sample reacts with the antibodies against Norovirus GI (strip A) and/or Norovirus GII (strip B), forming conjugates. The mixture moves upward on the membrane by capillary action. If the sample is Norovirus GI positive, antibodies present on the membrane (test line) capture the conjugate complex and a **red** line will be visible in the strip A, and if the sample is Norovirus GII positive, antibodies present on the membrane (test line) capture the conjugate complex and a **red** line will be visible in strip B. Although the sample is positive or negative, the mixture continues to move across the membranes and the **green** control line always appears (for both strips).

The presence of these **green** lines (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 expiration date.
- Do not use the test if its pouch is damaged.
- Specimens should be considered as potentially hazardous and handle in the same manner as an infectious agent. A new test must be used for each sample to avoid contaminations errors. Device for single use.
- Tests should be discarded in a proper biohazard container after testing.
- Reagents contain preservatives. Avoid any contact with the skin or mucous membrane. Consult safety data sheet, available on request.
- Components provided in the kit are approved for use with the **Vitassay Norovirus GI+GII**. Do not use any other commercial kit component.
- Follow Good Laboratory Practices, wear protective clothing, use disposal gloves, goggles and mask. Do not eat, drink or smoke in the working area.

STORAGE AND STABILITY

Store as packaged in the sealed pouch either at refrigerated or room temperature (2-30°C/35.6-86°F).

The test is stable until the expiration date printed on the sealed pouch.

The test must remain in the sealed pouch until use.

Do not freeze.

MATERIALS

MATERIAL PROVIDED	MATERIAL REQUIRED BUT NOT PROVIDED
<ul style="list-style-type: none"> 25 tests/kit Vitassay Norovirus GI+GII Instructions for use. 25 Vials with diluent for the sample dilution. 	<ul style="list-style-type: none"> Specimen collection container. Disposable gloves. Timer.

SPECIMEN COLLECTION

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

Samples can be stored in the refrigerator (2-8°C/35.6-46.4°F) for 1-2 days prior to testing. For longer storage, maximum 1 year, the specimen must be kept frozen at -20°C (-4°F). Samples must be brought to room temperature before testing.

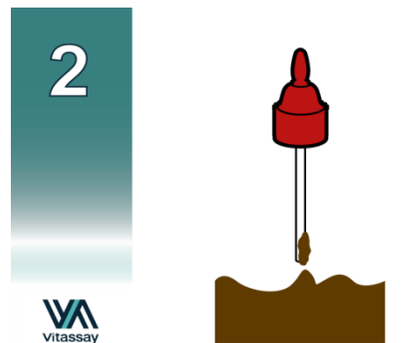
Homogenise stool samples as thoroughly as possible prior to preparation.

SPECIMEN PREPARATION

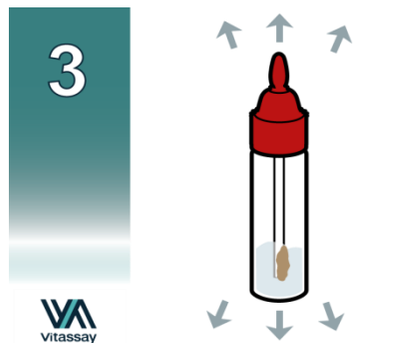
1. Remove the cap of the vial with diluent for the sample dilution (figure 1).
2. Use the stick to collect sufficient sample quantity (approx. 125mg). For solid stool, insert the stick in 4 different areas of the stool sample taken approx. 125mg, (figure 2), and add it into the vial with diluent for the sample dilution. For liquid stool, take 125 µL of the sample using a micropipette and transfer it into the vial with diluent for the sample dilution.
3. Close the tube with the diluent and stool sample. Shake vigorously the vial in order to assure good sample dispersion (figure 3).



Vial for sample dilution.



Insert the stick in 4 different areas of the stool.



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/59-86°F) prior to testing. Do not open pouches until the performance of the assay.

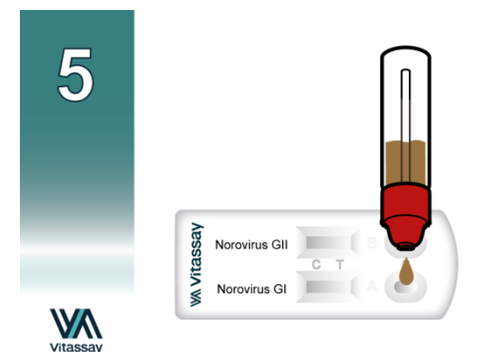
1. Shake the vial with the sample vigorously to obtain a good sample dilution.
2. Remove the **Vitassay Norovirus GI+GII** from its sealed bag just before using it.
3. Take the vial containing the diluted sample, cut the end of the cap (figure 4) and dispense 4 drops in the circular window marked with the letter S for the strip A – Norovirus GI (figure 5) and 4 drops, using the same vial, in the circular window marked with the letter S for the strip B – Norovirus GII (figure 6).
4. Read the results at **10 minutes**. Do not read the results later than 10 minutes.

If the test does not run due to solid particles, stir the sample added in the sample window with the stick. If it does not work,

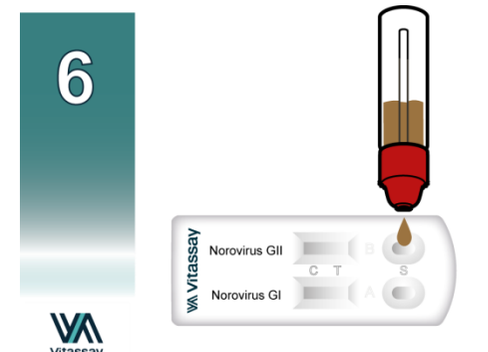
dispense a drop of diluent until seeing the liquid running through the reaction zone.



Cut the end of the cap.

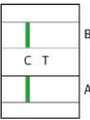
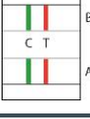
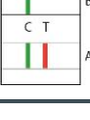
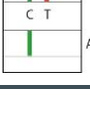


Dispense 4 drops in the circular window marked with the letter S to the strip A – Norovirus GI.



Dispense 4 drops in the circular window marked with the letter S to the strip B – Norovirus GII.

INTERPRETATION OF THE RESULTS

RESULTS	Strip A Norovirus GI	Strip B Norovirus GII	INTERPRETATION
	Negative GREEN	Negative GREEN	There is no Norovirus GI or Norovirus GII presence. No infection caused by Norovirus GI or Norovirus GII.
	Positive GREEN-RED	Positive GREEN-RED	There is Norovirus GI and Norovirus GII presence. Infection caused by Norovirus GI and Norovirus GII.
	Positive GREEN-RED	Negative GREEN	There is Norovirus GI presence. Infection caused by Norovirus GI.
	Negative GREEN	Positive GREEN-RED	There is Norovirus GII presence. Infection caused by norovirus GII.
ANY OTHER RESULTS			Invalid result, we recommend repeating the assay using the sample with another test. Note: Wrong procedural techniques or deterioration of the reagents are the main reasons of control line failure. If the symptoms or situation still persist, discontinue using the test kit and contact your local distributor.

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

QUALITY CONTROL

Internal procedural controls are included in **Vitassay Norovirus GI+GII**. Green lines appearing in the in the results window are internal controls, which confirm sufficient specimen volume and correct procedural technique.

LIMITATIONS

- Vitassay Norovirus GI+GII** 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 Norovirus GI+GII** depends on the quality of the sample; Proper fecal specimens must be obtained.
- Positive results determine the presence of Norovirus GI and/or Norovirus GII in human stool samples. A positive result should be followed up with additional laboratory techniques (biochemical methods or by PCR) to confirm the results. A confirmed infection should only be made by a physician after the evaluation of all clinical and laboratory findings 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 values. If symptoms or situation still persist, a Norovirus determination should be carried out on a sample from other technique (for example PCR).

EXPECTED VALUES

The majority of norovirus outbreaks occur in health care settings (including long-term care facilities and hospitals), where the virus is predominantly spread from person to person. In addition, norovirus have also been identified in over 58% of the reported foodborne outbreaks in which an etiologic agent was determined. In the most recent disease burden estimates in the United States, norovirus causes 570 to 800 deaths, 56000 to 71000 hospitalizations, 400000 emergency room visits, and 1.7 to 1.9 million outpatient visits annually.

In countries that belong to temperate latitudes, most infections occur in the fall and winter and at least 70% of outbreaks are reported in semiclosed communities such as long-term-care facilities, schools, hospitals, and cruise ships.

PERFORMANCE CHARACTERISTICS

Clinical sensitivity and specificity

An evaluation was performed using **Vitassay Norovirus GI+GII** and these results were compared with a commercial test (Simple Norovirus, Operon) and confirmed by PCR.

Results were as follows:

		Simple Norovirus		
		Positive	Negative	Total
Vitassay Norovirus GI+GII	Positive	2	0	2
	Negative	0	48	48
	Total	2	48	50

Vitassay Norovirus GI+GII (norovirus GI) vs Simple Norovirus			
Sensitivity	Specificity	PPV	NPV
>99%	>99%	>99%	>99%

		PCR		
		Positive	Negative	Total
Vitassay Norovirus GI+GII	Positive	2	0	2
	Negative	0	48	48
	Total	2	48	50

Vitassay Norovirus GI+GII (norovirus GI) vs PCR			
Sensitivity	Specificity	PPV	NPV
>99%	>99%	>99%	>99%

		Simple Norovirus		
		Positive	Negative	Total
Vitassay Norovirus GI+GII	Positive	10	0	10
	Negative	0	48	48
	Total	10	48	58

Vitassay Norovirus GI+GII (norovirus GII) vs Simple Norovirus			
Sensitivity	Specificity	PPV	NPV
>99%	>99%	>99%	>99%

		PCR		
		Positive	Negative	Total
Vitassay Norovirus GI+GII	Positive	8	0	8
	Negative	2	48	50
	Total	10	48	58

Vitassay Norovirus GI+GII (norovirus GII) vs PCR			
Sensitivity	Specificity	PPV	NPV
80%	>99%	>99%	96%

The results showed that **Vitassay Norovirus GI+GII** has a high sensitivity and specificity to detect Norovirus GI and Norovirus GII.

Cross reactivity

No cross reactivity was detected against other gastrointestinal pathogens that are occasionally present in feces:

<i>Adenovirus</i>	<i>Hepatitis A</i>	<i>Salmonella paratyphi</i>
<i>Astrovirus</i>	<i>Giardia Lamblia</i>	<i>Salmonella typhimurium</i>
<i>Campylobacter coli</i>	<i>Helicobacter pylori</i>	<i>Salmonella typhi</i>
<i>Campylobacter jejuni</i>	<i>Listeria monocytogenes</i>	<i>Shigella boydii</i>
<i>Clostridium difficile</i>	<i>Norovirus GI (Strip B)</i>	<i>Shigella dysenteriae</i>
<i>Cryptosporidium parvum</i>	<i>Norovirus GII (Strip A)</i>	<i>Shigella flexneri</i>
<i>Enterovirus</i>	<i>Rotavirus</i>	<i>Shigella sonnei</i>
<i>Escherichia coli O111</i>	<i>RSV</i>	<i>Staphylococcus aureus</i>
<i>Escherichia coli O157:H7</i>	<i>Salmonella enteritidis</i>	<i>Yersinia enterocolitica</i>

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SYMBOLS FOR IVD COMPONENTS AND REAGENTS

IVD	in vitro diagnostic device		Keep dry
	Consult instructions for use		Temperature limitation
	Use by		Manufacturer
LOT	Batch code		Contains sufficient for <n> test
DIL	Sample diluent	REF	Catalogue number

