

Rapid-VIDITEST

Adenovirus Card

One step Adenovirus Card for the qualitative detection of Adenovirus in faeces.

Instruction manual

INTENDED USE:

The Rapid-VIDITEST Adenovirus Card is a rapid chromatographic immunoassay for the qualitative detection of Adenovirus antigens in human faecal specimens to aid in the diagnosis of Adenovirus infection.

INTRODUCTION:

Viral gastroenteritis is an infection caused by a variety of viruses that results in vomiting or diarrhea. Many different viruses can cause gastroenteritis, including rotaviruses, noroviruses, adenoviruses, sapoviruses, and astroviruses.

The main symptoms of viral gastroenteritis are watery diarrhea and vomiting. The affected person may also have headache, fever, and abdominal cramps ("stomach ache"). In general, the symptoms begin 1 to 2 days following infection with a virus that causes gastroenteritis and may last for 1 to 10 days, depending on which virus causes the illness. Some research studies have shown that the duration of the symptoms are approximately three to four days. Adenoviruses and astroviruses cause diarrhea mostly in young children, but older children and adults can also be affected.

PRINCIPLE:

The Rapid-VIDITEST Adenovirus Card is a qualitative lateral flow immunoassay for the detection of Adenovirus antigen in human faeces samples. The membrane is pre-coated with monoclonal antibodies against Adenovirus antigens on the test line region. During testing, the sample reacts with the particle coated with anti-Adenovirus antibodies which was predried on the test strip. The mixture moves upward on the membrane by capillary action. In the case of a positive result the specific antibodies present on the membrane will react with the mixture conjugate and generate a coloured line. A green coloured band always appears in the control line and serves as verification that sufficient volume was added, that proper flow was obtained and as an internal control for the reagents.

MATERALS PROVIDED:

- Rapid-VIDITEST Adenovirus Card tests
- Instructions for use
- Specimen collection vial with buffer

MATERIALS REQUIRED BUT NO PROVIDED:

- Specimen collection container
- Disposable gloves
- Timer

SPECIMEN COLLECTION AND PREPARATION:

Collect sufficient quantity of faeces (1-2 g or mL for liquid sample). Stool samples should be collected in clean and dry containers (no preservatives or transport media). The samples can be stored in the refrigerator $(2-4^{\circ}C/36-40^{\circ}F)$ for 1-2 days prior to testing. For longer storage the specimen must be kept frozen at $-20^{\circ}C/-4^{\circ}F$. In this case, the sample will be totally thawed, and brought to room temperature before testing.

PROCEDURES:

To process the collected stool samples:

Use a separate specimen collection vial/testing tube for each sample. Dispense exactly 1mL of the buffer into a vial. Introduce the swab or stick two times into the faecal specimen to pick up a little sample (100 mg) (1) and put into the testing tube or vial with buffer. Shake the testing tube or vial in order to assure good sample dispersion (2). For liquid stool samples, aspirate the faecal specimen with a dropper and add 100 μ L into the testing tube or vial with buffer.



Test Procedure:

Allow the tests, stool samples and buffer to reach to room temperature (15-30°C/59-86°F) prior to testing. Do not open the pouch until ready to perform the assay.

- 1. Remove the Rapid-VIDITEST Adenovirus Card from its sealed pouch and use it as soon as possible.
- 2. Shake the specimen collection vial to assure a good sample dispersion. Break off the cap of the vial (3).
- 3. Use a separate device for each sample. Dispense exactly 4 drops into the specimen well (S) (4). Start the timer.
- 4. Read the result at **10 minutes** after dispensing the sample.



INTERPRETATION OF RESULTS:



POSITIVE: Two lines appears across the central window in the result line region, a **blue** test line marked with the letter T and in the control line region, a **green** control line marked with the letter C.

NEGATIVE: Only one **green** band appears across the control line region marked with the letter C.

INVALID: A total absence of the green control coloured band regardless the appearance or not of the blue test line. Note: Insufficient specimen volume, incorrect procedural techniques or deterioration of the reagents are the most likely reasons for control line failure. Review the procedure and repeat the test with a new test. If the problem persists, discontinue using the test kit and contact you local distributor.

NOTES ON THE INTERPRETATION OF RESULTS:

The intensity of the blue coloured band in the result line region (T) will vary depending on the concentration of viral antigens in the specimen. However, neither the quantitative value, nor the rate of increase in antigens can be determined by this qualitative test.

QUALITY CONTROL:

Internal procedural controls are included in the test:

A green line appearing in the control line region (C). It confirms sufficient specimen volume and correct procedural technique.

LIMITATIONS:

- 1. Rapid-VIDITEST Adenovirus Card will only indicate the presence of Adenovirus in the specimen (qualitative detection) and should be used for the detection of Adenovirus antigens in faeces specimens only. Neither the quantitative value nor the rate of increase in Adenovirus antigens concentration can be determined by this test.
- 2. An excess of sample could cause wrong results (brown bands appear). Dilute the sample with the buffer and repeat the test.
- 3. Some stool samples can decrease the intensity of the control line.
- 4. If the test result is negative and clinical symptoms persist, additional testing using other clinical methods is recommended. A negative result does not at any time preclude the possibility of Adenovirus infection.
- 5. This test provides a presumptive diagnosis of Adenovirus infections. All results must be interpreted together with other clinical information and laboratory findings available to the physician.

EXPECTED VALUES:

Adenoviruses cause diarrhea mostly in young children, but older children and adults can also be affected. Adenovirus infections occur throughout the year.

PERFORMANCE CHARACTERISTICS:

Sensitivity and specificity

An evaluation was conducted comparing the results obtained using the Rapid-VIDITEST Adenovirus Card to a commercial available Adenovirus ELISA assay.

Rapid-VIDITEST Adenovirus Card was highly specific (>99%) and also highly sensitive (>99%) compared with the results of that ELISA assay.

Cross-reactivity

It was performed an evaluation to determine the cross reactivity of Rapid-VIDITEST Adenovirus Card. There is not cross reactivity with common gastrointestinal pathogens, other organisms and substances occasionally present in feces: Astrovirus, Rotavirus, *Escherichia coli, Campylobacter, Giardia lamblia*, human Hemoglobin.

STORAGE AND STABILITY:

Store as packaged in the sealed pouch either at refrigerated or room temperature $(2-30^{\circ}C/36-86^{\circ}F)$. The test is stable through the expiration date printed on the sealed pouch. The test must remain in the sealed pouch until use. Do not freeze.

PRECAUTIONS:

- For professional in vitro diagnostic use only.
- Do not use after expiration date.
- The test should remain in the sealed pouch until use.
- Do not use the test if pouch is damaged.
- Follow Good Laboratory Practices, wear protective clothing, use disposal gloves, do not eat, drink or smoke in the area.
- All the specimens should be considered potentially hazardous and handled in the same manner as an infectious agent.
- The test should be discarded in a proper biohazard container after testing.
- The test must be carried out within 2 hours of opening the sealed bag.

REFERENCES:

1. GUILLERMO BERNAOLA, WALTER LUQUE. et al., "Fisiopatología de las Infecciones por Adenovirus", Paediatrica Asociación de Médicos Residentes del Instituto de Salud del Niño Oct. 2001 - Mar. 2002 Volumen 4, Nº 2 Págs. 41 - 47.

SYMBOLS FOR IVD COMPONENTS AND REAGENTS:

IVD	In vitro diagnostic device	LOT	Batch code
\sum	Use by		Manufacturer



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