RIDA®QUICK
Cryptosporidium/Giardia Combi

Article no: N1123
1. Intended use

For in vitro diagnostic use. The RIDA®QUICK Cryptosporidium/Giardia Combi is a quick immunochromatographic test for the qualitative determination of Cryptosporidium parvum and/or Giardia lamblia in stool samples.

2. Summary and explanation of the test

**Giardia lamblia** is an intestinal flagellate. The morphologically characteristic Trophozoites only survive for a short time outside the host organism. Transmission is affected via the highly infectious cysts. Because it is spread world-wide, Giardia lamblia has become an important cause of chronic diarrhoeas, particularly with the problems in travel medicine. The infection occurs after the ingestion of cysts in contaminated food and water. In communal facilities with inadequate hygiene, the infection usually occurs via the faecal-oral route from person to person. This mode of transmission is particularly common among children and in kindergartens, as well as among male homosexuals and inmates. The infection can also be passed on from the children to their parents. Unlike infants, older children who are infected can be free of symptoms. Nevertheless, they excrete the cysts and can infect other humans. The symptoms of Giardiasis (Lambliasis) are acute or chronic diarrhoea. The incubation time is between 3 and 42 days. The method most frequently used to diagnose Giardiasis in the past has been the detection by microscopy of cysts in the stool, which requires the services of experienced personnel. As well as this, the investigations have to be carried out over a long period of time, because the excretion of cysts is subject to strong fluctuations.

**Cryptosporidium parvum** is a parasite which is far more common in animals and occurs as an important pathogenic organism in domestic animals and in calves in particular. However, infections in humans are observed in many countries now more frequently than was accepted earlier. In tropical developing countries, the parasite is often endemic and causes diarrhoea epidemics among children. With immunocompetent patients, the disease manifests itself as self-healing gastroenteritis. The diarrhoea lasts between 3 and 10 days and may be accompanied by fever and gastrointestinal symptoms such as nausea and pain, which resembles those of Giardiasis (Lambliasis). The symptoms and effects are substantially more serious with immunoincompetent patients, where diarrhoeas persist and are very severe. The infection can be transmitted from animal to humans via contaminated water and from human to human. Members of communal facilities, children in kindergartens and the high-risk groups, homosexual men and patients infected with HIV, are particularly at risk. In the past, the methods most frequently used for the diagnosis of cryptosporidiosis were the microscopic detection of Oocysts in the stool or the microscopic examination of small intestine biopsy samples which required the services of experienced personnel.
An important alternative method to microscopy for the detection of Giardia lamblia and Cryptosporidium parvum is the quick immunochromatographic test described in the following which, because it uses monoclonal antibodies, is equivalent to the microscopy investigation procedures in terms of sensitivity and specificity. The test is quick and simple to perform and does not require specially trained microbiologists to carry it out.

3. Test principle

This quick test is a single-step immunochromatographic lateral-flow test, where specific antibodies which are directed against both parasites attach themselves to red (Giardia specific) or blue (Cryptosporidia specific) latex particles. Other specific antibodies against the two pathogens are firmly bound to the membrane. The stool sample is first suspended in the extraction buffer and then precipitated. An aliquot portion of the clear supernatant of the sample is placed on the test strip. The sample then passes, with the coloured latex particles to which antigen are attached if the test is positive, through the membrane and is bonded to the specific catch band. A blue and/or red band appears, depending on the antigens present in the sample.

4. Reagents provided

There are enough reagents in the pack for 20 determinations.

- **Cassette**: 20 det. 20 individually packed test cassettes
- **Diluent**: 26 ml Extraction buffer, ready for use; contains 0.1% sodium azide
- **Pipet**: 25 ea.. Bag with 25 disposable pipettes

5. Storage instructions

The pack can be stored at 2 – 30 °C and can be used until the printed expiry date. After the expiry date, the quality guarantee is no longer valid. Likewise, the usability of the cassettes cannot be guaranteed once the external packaging of the individual cassette has been damaged.
6. **Materials required but not provided**

- Test tubes for stool suspension
- Vortex mixer (optional)
- Micropipette (200 µl - 1000 µl)
- Waste container containing 0.5 % sodium hypochlorite solution

7. **Precautions for users**

For *in vitro* diagnostic use only. This test must only be carried out by trained laboratory personnel. The guidelines for working in medical laboratories must be followed and the instructions for carrying out the test must be strictly adhered to.

The sample dilution buffer contains sodium azide as a preservative. This substance must not be allowed to come into contact with the skin or mucous membrane.

Samples or reagents must not be pipetted by mouth and contact with injured skin or mucous membranes must be prevented. When handling the samples, wear disposable gloves and when the test is finished, wash your hands. Do not smoke, eat or drink in areas where samples or test reagents are being used.

All reagents and materials which come into contact with potentially infectious samples must be treated exactly like the samples themselves with suitable disinfectants (e.g. sodium hypochlorite) or autoclaved for at least one hour at 121 °C.

8. **Specimen collection and storage**

Stool samples must be collected in clean containers without any additives and stored at 2 – 8 °C before beginning of test. If stored for more than 3 days, the sample must be frozen at -20 °C. In this case, the sample must be completely thawed out and brought to room temperature before testing begins. Multiple freezing and thawing of the sample must be avoided.

If rectal swabs have to be used, make sure that sufficient stool material (approx. 50 mg) is collected to carry out the test.
9. Test procedure

9.1. General information

Before using the samples, the extraction buffer and the test cassettes must be brought to room temperature (20 – 25 °C). The test cassettes must only be removed from the external packaging shortly before they are used. Once used, the cassettes must not be used again. The test must not be carried out in direct sunlight.

Do not pour reagents back into vials as reagent contamination may occur.

9.2. Preparing the samples

Place 1 ml Extraction Buffer Diluent in the test tubes indicated. With the liquid stool sample, pipette 100 µl of the sample with a disposable pipette Pipet (up to just above the second thickening) and suspend it in the buffer placed in the tube beforehand. With solid stool samples, suspend 50 mg in the buffer. The sample must then be well homogenized. This can be achieved either by repeated suction and ejection of the suspension using the disposable pipette Pipet or, alternatively, by mixing on a vortex mixer. Afterwards, allow the homogeneous suspension to settle for at least 3 minutes until a clear supernatant is formed.

9.3. Testing the sample

When removed from the external packing, first lay the test cassette on a level mat. After this, pipette 200 µl of the clear supernatant of the stool suspension with a micropipette or 4 drops with a disposable pipette Pipet into the round opening of the test cassette. Makes sure that the liquid flows through the membrane unimpeded. Any particles pipetted at the same time can cause an obstruction and must be removed beforehand. The test result can be read off after 5 minutes.

10. Quality control – indications of reagent expiry

The test must only be evaluated if the test cassette is intact before the sample suspension is pipetted in and no colour changes or bands are visible on the membrane. In addition to this, at least the green control band must be visible after the test incubation. If this does not appear, the following must be checked before repeating the test:

- Expiry date of the test cassettes and the extraction buffer being used
- Correct test procedure
- Contamination of the extraction buffer

After this, if the control band is still not visible after repeating the test with a new test cassette, please contact the manufacturer.
11. Evaluation and interpretation

A maximum of three bands should appear in the following order, as seen from the sample-absorption site: one blue ($T_1 = \text{Test band 1}$), one red ($T_2 = \text{Test band 2}$) and one green ($C = \text{Control band}$) band. If the green control band is missing, the test is invalid and cannot be evaluated!

The following interpretations are possible:

- **Cryptosporidium positive**: blue and green bands are visible.
- **Giardia positive**: red and green bands are visible.
- **Cryptosporidium and Giardia positive**: blue, red and green bands are visible.
- **Negative**: only the green band is visible.
- **Not valid**: no visible band or a combination other than the one described above or other changes in band colour. Likewise, changes in band colour which only appear after 10 minutes or later are also without any diagnostic value and must not be used for evaluation.

12. Limitations of the method

The RIDA®QUICK Cryptosporidium/Giardia Combi test detects antigens of Cryptosporidium parvum and/or Giardia lamblia in stool samples. The test cannot be used to derive a relationship between the intensity of the specific visible bands and the occurrence or severity of clinical symptoms. The results obtained must always be interpreted in combination with the clinical picture.

A positive result does not rule out the presence of another infectious pathogen.

A negative result does not necessarily mean that there is no Cryptosporidia or Lamblien infection. This can be caused by intermittent excretion of the pathogen or by the quantity of antigens in the sample being too small. If the patient is anaemic or is suspected as being infected by the pathogens being looked for, another stool sample should be tested after four weeks.

An excess of stool sample can cause brownish bands to appear instead of the specifically coloured bands. These brownish bands do not have any diagnostic value. In such cases, it will be necessary to repeat the test with a smaller stool quantity or dilute the suspension already prepared further (clear supernatant after precipitation) in order to clarify whether the pathogens being looked for are in the sample and have been masked by too much stool matrix.
13. Performance characteristics

13.1. Clinical comparison study

In a routine laboratory, a comparative investigation was carried out between the RIDA®QUICK Cryptosporidium/Giardia Combi method and the established microscopy method on both frozen and fresh stool samples using 70 stool samples in total (15 Cryptosporidium positive and 15 Giardia lamblia positive as well as 40 negative stool samples). The results are listed in Tables 1 and 2.

Table 1: Comparison of the RIDA®QUICK Cryptosporidium/Giardia Combi with microscopy with exclusive evaluation of the Cryptosporidia-specific band

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Sensitivity: 93.8 %  pos. prognosis value: 100.0 %  Specificity: 100.0 %  neg. prognosis value: 97.5 %

Table 2: Comparison of the RIDA®QUICK Cryptosporidium/Giardia Combi with microscopy with exclusive evaluation of the Giardia-specific band

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Sensitivity: 100.0 %  pos. prognosis value: 88.2 %  Specificity: 95.2 %  neg. prognosis value: 100.0 %
13.2. Cross reactivity

None that led to a cross reaction in the following specified intestine parasites in the RIDA®QUICK Cryptosporidium/Giardia Combi test:

Entamoeba coli
Blastocystis hominis
Jodamoeba buetschlii
Chilomastix mesnili
Endolimax nana
Eggs of Taenia spp.
References