

RIDA® GENE ETEC/EIEC

REF PG2225



1. Intended use

For *in vitro* diagnostic use. RIDA[®]GENE ETEC/EIEC is a multiplex real-time PCR for the direct, qualitative detection and differentiation of the virulence-factor genes of ETEC and EIEC/*Shigella* spp. in human stool samples and cultures.^{1,2} RIDA[®]GENE ETEC/EIEC multiplex real-time PCR is intended for use as an aid in diagnosis of gastroenteritis caused by pathogenic *Escherichia coli* and *Shigella* spp., respectively.

2. Summary and explanation of the test

Escherichia coli (*E. coli*) are gram negative, facultatively anaerobic rod bacteria, which move by peritrichal flagellation and belong to the Enterobacteriaceae family. *E. coli* are part of the normal intestinal flora of humans and many farm animals and are generally nonpathogenic. Some *E. coli* strains are pathogenic to humans through the acquisition of certain virulence factors (e.g. genes for toxins).

The six known intestinal pathogenic *E. coli*: enterohemorrhagic *E. coli* (EHEC), enteropathogenic *E. coli* (EPEC), enterotoxigenic *E. coli* (ETEC), enteroinvasive *E. coli* (EIEC), EAEC enteroaggregative *E. coli* und diffusely adherent *E. coli* (DAEC) can be differentiated by the virulence factors.¹

Enterotoxigenic *E. coli* (ETEC) are important intestinal pathogenic *E. coli*. ETEC is the most common cause of traveller's diarrhea that affects persons travelling to developing countries. 30 - 60 % of all traveller's diarrhea cases are caused by ETEC. 11 – 16 % of returning travellers from developing countries are infected by ETEC. Among children below 5 years of age in developing countries, ETEC accounts for about 210 million cases of diarrhea and about 380,000 deaths annually.²

ETEC has two important diagnostic virulence factors. They are capable to produce heat stable (ST) and/or heat labile (LT) enterotoxins. The clinical symptoms which are caused by ETEC range from mild to acute watery diarrhea, which fade without any specific therapy. Life-threatening cases of ETEC diarrhea occur mostly in weanling infants in the developing countries. Sources of infection are contaminated water and food, whereas infective chains from human to human can be excluded so far.³

Enteroinvasive *E. coli* (EIEC) are responsible for Shigellose-like disease in developing countries and among travellers to these less developed regions. EIEC strains are biochemically and genetically related to *Shigella* spp. The pathogenic features of EIEC and *Shigella* spp. are based on plasmid-mediated capability to invade the colonic epithelium for destruction.³ By the detection of the ipaH gene (invasion plasmid antigen H gene) EIEC/*Shigella* spp. can be differentiated from ETEC. A differentiation between EIEC and *Shigella* spp. is not recommended, because both have the same clinical relevance. Shigellose caused by EIEC is characterized by abdominal pain and watery diarrhea, sometimes containing blood. Sources of infection are contaminated water, food and infective chains from human to human.⁴

3. Test principle

RIDA[®]GENE ETEC/EIEC is a multiplex real-time PCR for the direct, qualitative detection and differentiation of the virulence-factor genes of ETEC and EIEC/*Shigella* spp. in human stool samples and cultures.

After DNA-isolation, amplification of the gene fragments specific for the heat-labile enterotoxin LT (elt), the heat-stable enterotoxin ST (estA) and the ipaH-gene (if present) occurs. The amplified targets are detected with hydrolysis probes, which are labeled at one end with a quencher and at the other end with a fluorescent reporter dye (fluorophore). In the presence of a target the probes hybridize to the amplicons. During the extension step the **Taq-Polymerase** breaks the reporter-quencher proximity. The reporter emits a fluorescent signal which is detected by the optical unit of a real-time PCR instrument. The fluorescence signal increases with the amount of formed amplicons. The RIDA[®]GENE ETEC/EIEC assay contains an **Internal Control DNA** (ICD) as an internal control of sample preparation procedure and to determine possible PCR-inhibition.

4. Reagents provided

Tab. 1: Reagents provided (Reagents provided in the kit are sufficient for 100 determinations).

| Kit Code | Reagent | Amount | | Lid Color |
|----------|----------------------|--------|---------|-----------|
| 1 | Reaction Mix | 2x | 1050 µl | yellow |
| 2 | Taq-Polymerase | 1x | 80 µl | red |
| D | Internal Control DNA | 2x | 1700 µl | orange |
| N | No Template Control | 1x | 450 µl | white |
| P | Positive Control | 1x | 200 µl | blue |

5. Storage instructions

- Protect all reagents from light and store at -20 °C. All reagents can be used until the expiration date. After expiry the quality guarantee is no longer valid.
- Carefully thaw reagents before using (e.g. in a refrigerator at 2 - 8 °C).
- Reagents can sustain up to **20 freeze/thaw cycles** without influencing the assay performance (e.g. after the first thawing separate it in aliquots and freeze immediately).
- During PCR preparation all the reagents should be stored cold in an appropriate way (2 - 8 °C).

6. Additional necessary reagents and necessary equipment

The RIDA[®] GENE ETEC/EIEC multiplex real-time PCR assay is suitable for use with following extraction platforms and real-time PCR instruments:

Tab. 2: Validated equipment

| Extraction platforms | |
|---------------------------|--|
| R-Biopharm | RIDA [®] Xtract |
| Promega | Maxwell [®] RSC |
| bioMérieux | NucliSENS [®] easyMag [®] |
| Real-time PCR instruments | |
| Roche | LightCycler [®] 480II, LightCycler [®] 480 z |
| Agilent Technologies | Mx3005P |
| Applied Biosystems | ABI 7500 |
| Bio-Rad | CFX96 [™] |
| QIAGEN | Rotor-Gene Q |

Note: Only use 0.1 ml tubes on the Rotor-Gene Q (QIAGEN).

If you want to use other extraction platforms or real-time PCR instruments please contact R-Biopharm at pcr@r-biopharm.de.

- RIDA[®] GENE Color Compensation Kit IV (PG0004) for use with the LightCycler[®] 480II and the LightCycler[®] 480 z
- Real-time PCR consumables (plates, tubes, foil)
- Centrifuge with a rotor for the reaction vials
- Vortexer
- Pipettes (0.5 - 20 µl, 20 - 200 µl, 100 - 1000 µl)
- Filter tips
- Powder-free disposal gloves
- PCR water (BioScience grade, nuclease-free)

7. Precautions for users

For *in-vitro* diagnostic use.

This test must only be carried out by trained laboratory personnel. The guidelines for working in medical laboratories have to be followed. The instruction manual for the test procedure has to be followed. Do not pipet samples or reagents by mouth. Avoid contact with bruised skin or mucosal membranes. During handling reagents or samples, wear appropriate safety clothing (appropriate gloves, lab coat, safety goggles) and wash your hands after finishing the test procedure. Do not smoke, eat or drink in areas where samples or reagents are being used.

- Extraction, PCR preparation and the PCR run should be separated in different rooms to avoid cross-contaminations.
- Samples must be treated as potentially infectious as well as all reagents and materials being exposed to the samples and have to be handled according to the national safety regulations.
- Do not use the kit after the expiration date.

All reagents and materials used have to be disposed properly after use. Please refer to the relevant national regulations for disposal.

For more details see Safety Data Sheets (SDS) at www.r-biopharm.com.

8. Collection and storage of samples

8.1 Sample preparation from stool samples

For DNA isolation of human stool samples, use a commercially available DNA isolation kit (e.g. RIDA[®] Xtract (R-Biopharm)) or DNA extraction system (e.g. Maxwell[®] RSC (Promega)). Extract DNA according to the manufacturer's instructions.

We recommend to dilute the stool samples before extraction 1:3 with water. Vortex the diluted stool sample intensely and centrifuge at 1000 x g for 30 sec. Use from the supernatant the appropriate volume according to the manufacturer's instruction.

The RIDA[®]GENE ETEC/EIEC assay contains an **Internal Control DNA** that detects PCR inhibition, monitors reagent integrity and confirms that nucleic acid extraction was sufficient. The **Internal Control DNA** can either be used as PCR inhibition control or as extraction control for the sample preparation procedure and as PCR inhibition control.

If the **Internal Control DNA** is used only as a PCR inhibition control, 1 µl of the **Internal Control DNA** should be added to the Master-Mix (see Tab.4).

If the **Internal Control DNA** is used as an extraction control for the sample preparation procedure **and** as PCR inhibition control, 20 µl of the **Internal Control DNA** has to be added during extraction procedure.

The **Internal Control DNA** should always be added to the specimen-lysis buffer mixture and must **not** be added directly to the specimen. We also recommend to add

1 µl of the **Internal Control DNA** to the negative control and positive control PCR Mix.

8.2 Sample preparation from cultures

For DNA isolation from culture the following procedure is recommended: Add 1 ml PCR water into a preparation tube. Collect colonies with an inoculation loop and suspend them in the prepared PCR water. Cut or break the inoculation loop stem. Cap the preparation tube tightly and vortex strongly for 60 seconds. Heat and shake the preparation tube at 95 °C for 10 min in a heating block. Centrifuge for 1 min at 13.000 x g and apply the supernatant as sample.

Note: Repeat the centrifugation step in case of strong turbidity (if necessary).

The RIDA[®]GENE ETEC/EIEC assay contains an **Internal Control DNA** that detects PCR inhibition, monitors reagent integrity and confirms that nucleic acid extraction was sufficient. The **Internal Control DNA** can either be used as PCR inhibition control or as extraction control for the sample preparation procedure and as PCR inhibition control.

If the **Internal Control DNA** is used only as a PCR inhibition control, 1 µl of the **Internal Control DNA** should be added to the Master-Mix (see Tab.4).

If the **Internal Control DNA** is used as an extraction control for the sample preparation procedure **and** as PCR inhibition control, 20 µl of the **Internal Control DNA** has to be added during extraction procedure.

The **Internal Control DNA** should always be added to the specimen-PCR water mixture and must **not** be added directly to the specimen. We also recommend to add 1 µl of the **Internal Control DNA** to the negative control and positive control PCR Mix.

9. Test procedure

9.1 Master-Mix preparation

Calculate the total number of PCR reactions (sample and control reactions) needed. One positive control and one negative control must be included in each assay run.

We recommend calculating an additional volume of 10% to compensate imprecise pipetting (see Tab. 3, Tab. 4). Thaw, mix gently and briefly centrifuge the **Reaction Mix**, the **Taq-Polymerase**, the **Positive Control**, the **No Template Control** and the **Internal Control DNA** before using. Keep reagents appropriately cold during working step (2 - 8 °C).

Tab. 3: Calculation and pipetting example for 10 reactions of the Master-Mix (ICD as extraction and PCR inhibition control)

| Kit code | Master-Mix components | Volume per reaction | 10 reactions (10 % extra) |
|----------|-----------------------|---------------------|---------------------------|
| 1 | Reaction Mix | 19.3 µl | 212.3 µl |
| 2 | Taq-Polymerase | 0.7 µl | 7.7 µl |
| | Total | 20 µl | 220 µl |

Mix the components of the Master-Mix gently and briefly spin down.

Tab. 4: Calculation and pipetting example for 10 reactions of the Master-Mix (ICD only as PCR inhibition control)

| Kit code | Master-Mix components | Volume per reaction | 10 reactions (10 % extra) |
|----------|-----------------------|---------------------|---------------------------|
| 1 | Reaction Mix | 19.3 µl | 212.3 µl |
| 2 | Taq-Polymerase | 0.7 µl | 7.7 µl |
| D | Internal Control DNA | 1.0 µl | 11 µl |
| | Total | 21.0 µl | 231.0 µl |

Mix the components of the Master-Mix gently and briefly spin down.

9.2 Preparation of the PCR-Mix

Pipette 20 µl of the Master-Mix in each reaction vial (tube or plate).

Negative control: Add 5 µl **No Template Control** to the pre-pipetted Master-Mix.

Note: If the **Internal Control DNA** is used as extraction control for the sample preparation procedure and as PCR inhibition control, we recommend to add 1 µl of the **Internal Control DNA** to the PCR-Mix of the negative control.

Sample: Add 5 µl DNA extract to the pre-pipetted Master-Mix.

Positive control: Add 5 µl **Positive Control** to the pre-pipetted Master-Mix.

Note: If the **Internal Control DNA** is used as extraction control for the sample preparation procedure and as PCR inhibition control, we recommend to add 1 µl of the **Internal Control DNA** to the PCR-Mix of the positive control.

Cover tubes or plate. Spin down and place in the real-time PCR instrument. The PCR reaction should be started according to the PCR instrument set-up (Tab. 5, Tab. 6, Tab. 7, Tab. 8).

9.3 PCR instrument set-up

9.3.1 DNA real-time PCR profile

Tab. 5: DNA real-time PCR profile for LightCycler® series and Rotor-Gene Q

| | |
|---|---------------|
| Initial Denaturation | 1 min, 95 °C |
| Cycles | 45 Cycles |
| <u>PCR</u> Denaturation | 10 sec, 95 °C |
| Annealing/Extension | 15 sec, 60 °C |
| Temperature Transition Rate / Ramp Rate | Maximum |

Note: Annealing and Extension occur in the same step.

Tab. 6: DNA real-time PCR profile for Mx3005P, ABI 7500 and CFX96™

| | |
|---|---------------|
| Initial Denaturation | 1 min, 95 °C |
| Cycles | 45 Cycles |
| <u>PCR</u> Denaturation | 15 sec, 95 °C |
| Annealing/Extension | 30 sec, 60 °C |
| Temperature Transition Rate / Ramp Rate | Maximum |

Note: Annealing and Extension occur in the same step.

9.3.2 Universal real-time PCR profile

Note: The universal real-time PCR profile should only be used for DNA assays when combining RIDA® GENE DNA and RNA real-time PCR assays in one run.

Tab. 7: Universal real-time PCR profile for LightCycler® series

| | |
|---|---------------|
| <u>Reverse Transcription</u> | 10 min, 58 °C |
| Initial Denaturation | 1 min, 95 °C |
| Cycles | 45 Cycles |
| <u>PCR</u> Denaturation | 10 sec, 95 °C |
| Annealing/Extension | 15 sec, 60 °C |
| Temperature Transition Rate / Ramp Rate | Maximum |

Note: Annealing and Extension occur in the same step.

Tab. 8: Universal real-time PCR profile for Mx3005P, ABI 7500, Rotor-Gene Q and CFX96™

| | |
|---|---------------|
| <u>Reverse Transcription</u> | 10 min, 58 °C |
| Initial Denaturation | 1 min, 95 °C |
| Cycles | 45 Cycles |
| <u>PCR</u> Denaturation | 15 sec, 95 °C |
| Annealing/Extension | 30 sec, 60 °C |
| Temperature Transition Rate / Ramp Rate | Maximum |

Note: Annealing and Extension occur in the same step.

9.4 Detection channel set-up

Tab. 9: Selection of appropriate detection channels

| Real-time PCR instrument | Detection | Detection channel | Note |
|----------------------------|-------------|-------------------|---|
| Roche LightCycler® 480II | LT | 465/510 | RIDA® GENE Color Compensation Kit IV (PG0004) is required |
| | ICD | 533/580 | |
| | ipaH | 533/610 | |
| | ST | 618/660 | |
| Roche LightCycler® 480 z | LT | 465/510 | RIDA® GENE Color Compensation Kit IV (PG0004) is required |
| | ICD | 540/580 | |
| | ipaH | 540/610 | |
| | ST | 618/660 | |
| Agilent Techn. Mx3005P | LT | FAM | Check that reference dye is none |
| | ICD | HEX | |
| | ipaH | ROX | |
| | ST | Cy5 | |
| Applied Biosystem ABI 7500 | LT | FAM | Check that passive reference option ROX is none |
| | ICD | VIC | |
| | ipaH | ROX | |
| | ST | Cy5 | |
| Bio-Rad CFX96™ | LT | FAM | - |
| | ICD | VIC | |
| | ipaH | ROX | |
| | ST | Cy5 | |
| Qiagen Rotogene Q | LT | Green | The gain settings have to be set to 5, according to the default settings |
| | ICD | Yellow | |
| | ipaH | Orange | |
| | ST | Red | |

10. Quality control

The analysis of the samples is done by the software of the used real-time PCR instrument according to the manufacturer`s instructions. Positive control and negative control have to show correct results (see Table 10, Fig. 1, Fig. 2, Fig.3) in order to determine a valid run.

The Positive Control CD Toxin A/B has a concentration of 10^3 copies/ μ l. In each PCR run it is used in a total amount of 5×10^3 copies.

Tab. 10: For a valid run, the following conditions must be met:

| Sample | Assay result | ICD Ct | Target Ct |
|------------------|--------------|------------------|-----------------------------------|
| Positive control | Positive | NA ^{*1} | See Quality Assurance Certificate |
| Negative control | Negative | Ct > 20 | 0 |

^{*1} No Ct value is required for the ICR to make a positive call for the positive control.

If the positive control is not positive within the specified Ct range but the negative control is valid, prepare all new reactions including the controls.

If the negative control is not negative but the positive control is valid, prepare all new reactions including the controls.

If the required criteria are not met, following items have to be checked before repeating the test:

- Expiry of the used reagents
- Functionality of the used instrumentation
- Correct performance of the test procedure

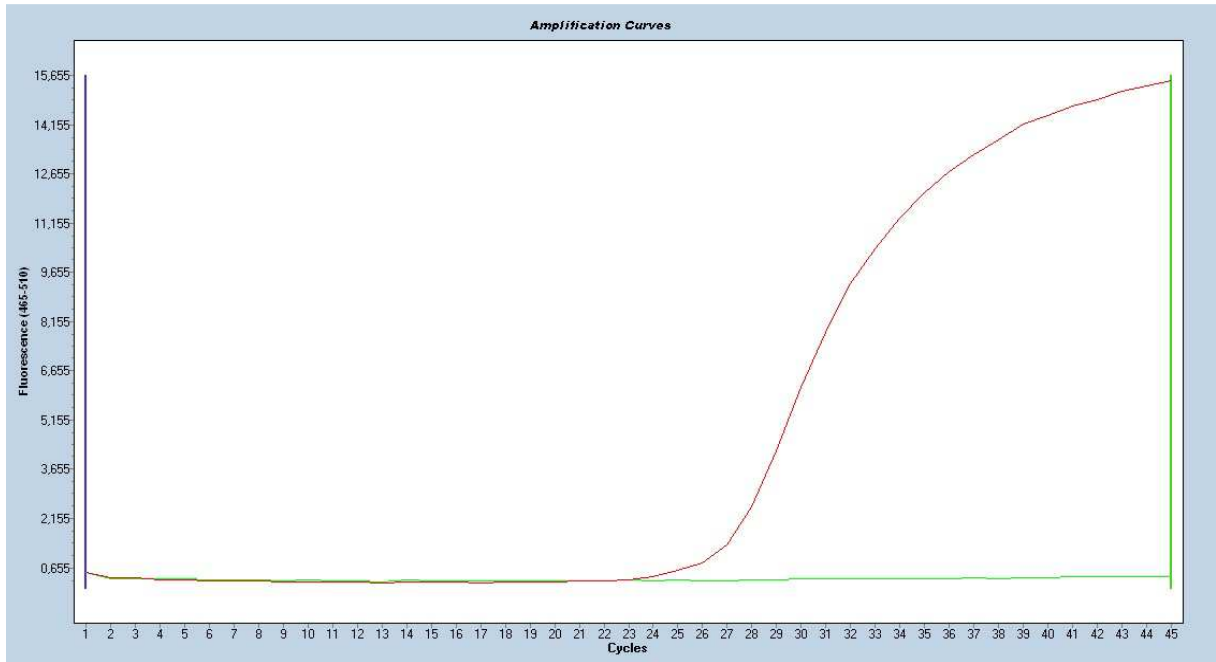


Fig. 1: Correct run of the positive and negative control (LT gene) on the LightCycler® 480II

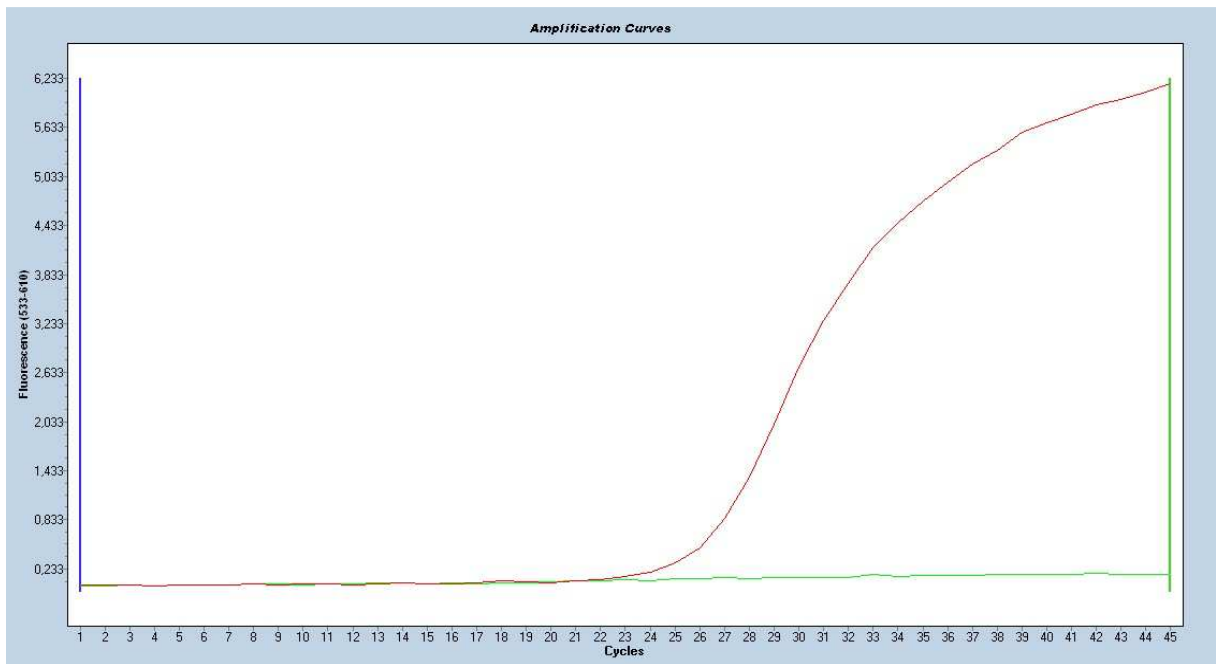


Fig. 2: Correct run of the positive and negative control (ipaH gene) on the LightCycler® 480II

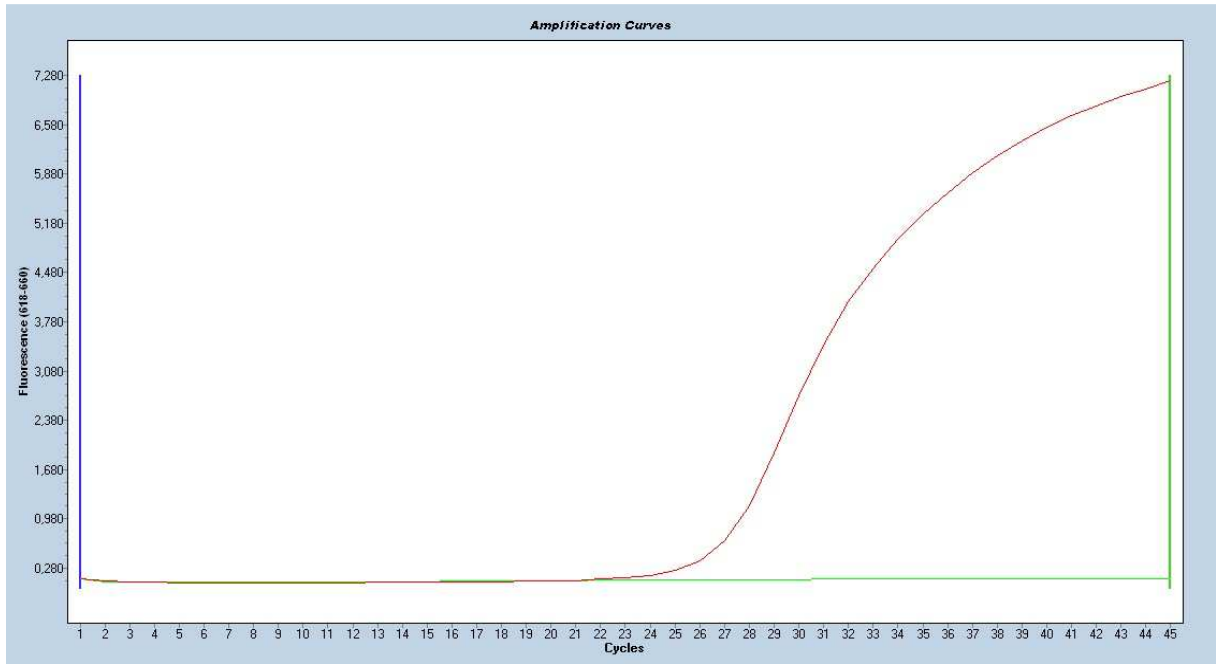


Fig. 3: Correct run of the positive and negative control (ST gene) on the LightCycler® 480II

11. Result interpretation

The result interpretation is done according to Table 11.

Tab. 11: Sample interpretation

| Virulence factor genes | | | | |
|------------------------|-----------------|-----------------|--------------------------|---|
| LT | ipaH | ST | ICD | Result |
| positive | negative | negative | positive/negative | ETEC detected |
| negative | positive | negative | positive/negative | EIEC/Shigella spp. detected |
| negative | negative | positive | positive/negative | ETEC detected |
| positive | positive | negative | positive/negative | ETEC und EIEC/Shigella spp. detected |
| positive | negative | positive | positive/negative | ETEC detected |
| negative | positive | positive | positive/negative | ETEC und EIEC/Shigella spp. detected |
| positive | positive | positive | positive/negative | ETEC und EIEC/Shigella spp. detected |
| negative | negative | negative | positive | Zielgene not detected |
| negative | negative | negative | negative | Invalid |

A sample is evaluated positive, if the sample DNA and the **Internal Control DNA** show an amplification signal in the detection system.

A sample is also evaluated positive, if the sample DNA shows an amplification signal but none for the **Internal Control DNA** in the detection system. The detection of the internal amplification control is not necessary because high concentrations of the amplicon can cause a weak or absent signal of the **Internal Control DNA**.

A sample is evaluated negative, if the sample DNA shows no amplification signal, but an amplification signal for the **Internal Control DNA** in the detection system. An inhibition of the PCR reaction can be excluded by the detection of the **Internal Control DNA**.

A sample is invalid, if the sample DNA and **Internal Control DNA** show no amplification signal in the detection system. The sample contains a PCR inhibitor or a

failure occurred in the extraction procedure. The extracted sample needs to be further diluted with PCR water (1:10) and re-amplified, or the isolation and purification of the sample has to be improved.

12. Limitations of the method

1. The result of molecular analysis should not lead to the diagnosis, but always be considered in the context of medical history and symptoms of the patient.
2. This assay is only validated for stool and culture samples.
3. Inappropriate specimen collection, transport, storage and processing or a pathogen load in the specimen below the analytical sensitivity can result in false negative results.
4. The presence of PCR inhibitors may cause invalid results.
5. Mutations or polymorphisms in primer or probe binding regions may affect detection of new variants resulting in a false negative result with the RIDA[®]GENE ETEC/EIEC assay.
6. As with all PCR based *in vitro* diagnostic tests, extremely low levels of target below the limit of detection (LoD) may be detected, but results may not be reproducible.
7. A positive test result does not necessarily indicate the presence of viable organisms. However, a positive result is indicative for the presence of the target genes (elt, estA or ipaH).

13. Performance characteristics

13.1 Analytical sensitivity

The RIDA[®]GENE ETEC/EIEC multiplex real-time PCR has a detection limit of ≥ 10 DNA copies per reaction.

The following figures 4, 5 and 6 show dilution series of the LT, ipaH and ST gene (each $10^5 - 10^1$ DNA copies per μl) on the LightCycler[®] 480II.

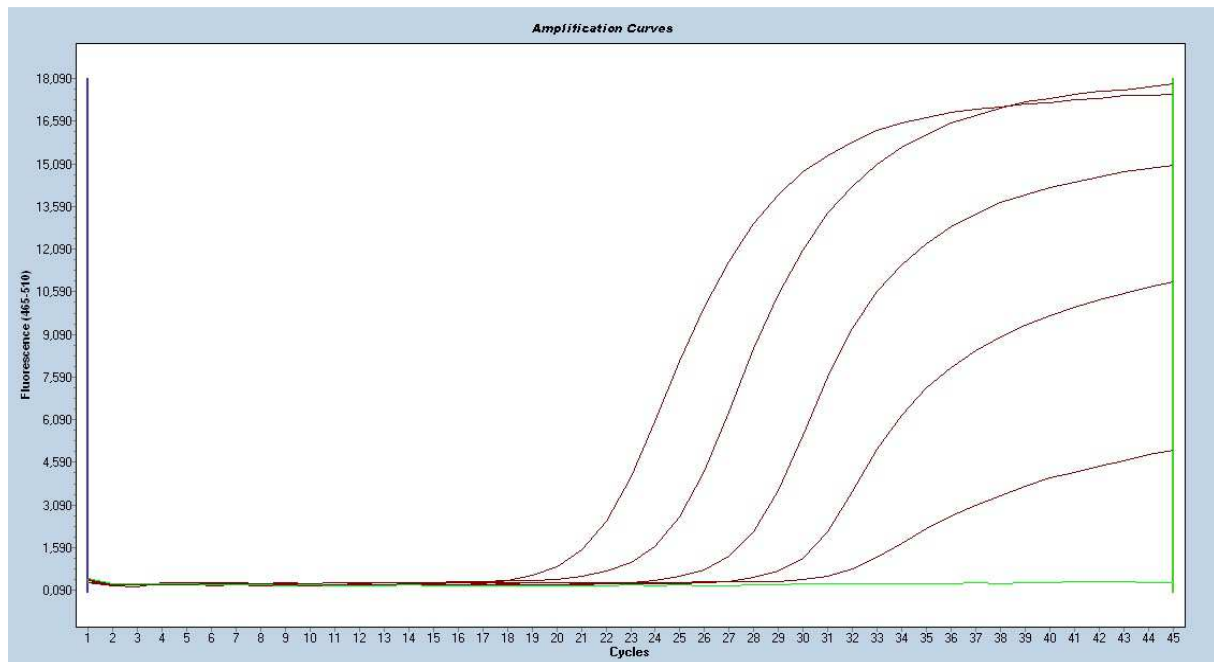


Fig. 4: Dilution series LT gene ($10^5 - 10^1$ DNA copies per μl) on the LightCycler[®] 480II

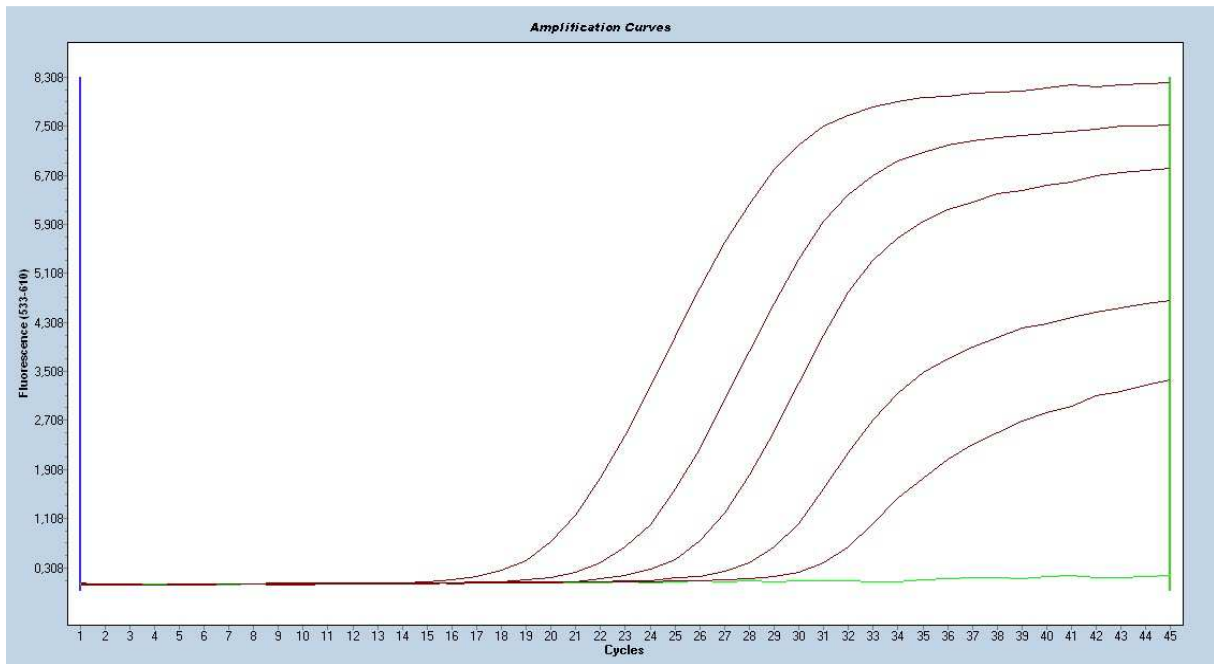


Fig. 5: Dilution series ipaH gene ($10^5 - 10^1$ DNA copies per μl) on the LightCycler[®] 480II

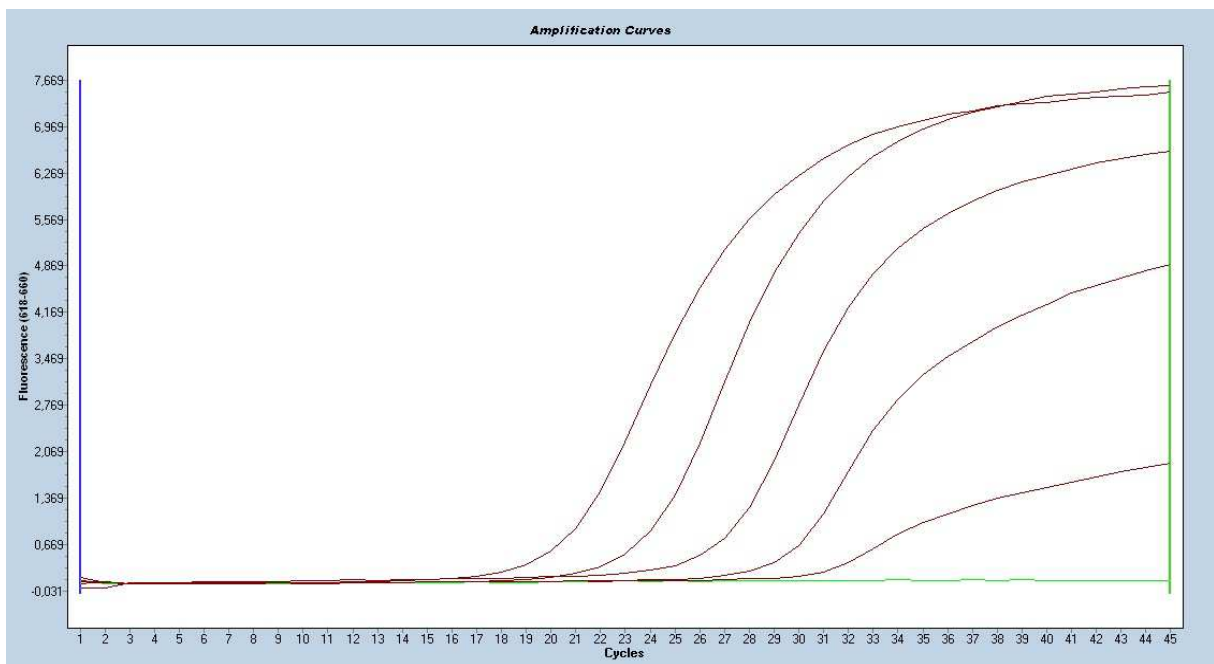


Fig. 6: Dilution series ST gene ($10^5 - 10^1$ DNA copies per μl) on the LightCycler[®] 480II

The detection limit of the whole procedure depends on the sample matrix, DNA extraction and DNA concentration.

13.2 Analytical specificity

The RIDA® GENE ETEC/EIEC multiplex real-time PCR is specific for ETEC and EIEC/*Shigella* spp. No cross-reaction could be detected for the following species (see Tab.12):

Tab. 12: Cross-reactivity testing










| | | | | | | | |
|--|---|--|---|---|---|-----------------------------------|---|
| Adenovirus 1, human, strain Adenoid 71 | - | <i>Campylobacter lari</i> subsp. <i>lari</i> | - | <i>Cryptosporidium parvum</i> | - | <i>Proteus vulgaris</i> | - |
| Adenovirus 7, human, strain Gomen | - | <i>Campylobacter upsaliensis</i> | - | <i>E. coli</i> (O157:H7) | - | <i>Pseudomonas aeruginosa</i> | - |
| Adenovirus 40, human, strain Dugan | - | <i>Candida albicans</i> | - | <i>E. coli</i> (O26:H-) | - | Rotavirus | - |
| Adenovirus 41, human, strain Tak | - | <i>Citrobacter freundii</i> | - | <i>Entamoeba histolytica</i> | - | <i>Salmonella enteritidis</i> | - |
| <i>Aeromonas hydrophila</i> | - | <i>Clostridium bifermentans</i> | - | <i>Enterobacter cloacae</i> | - | <i>Salmonella typhimurium</i> | - |
| <i>Arcobacter butzleri</i> | - | <i>Clostridium difficile</i> | - | <i>Enterococcus faecalis</i> | - | <i>Serratia liquefaciens</i> | - |
| Astrovirus | - | <i>Clostridium novyi</i> | - | <i>Giardia intestinalis</i> Portland 1 | - | <i>Shigella flexneri</i> | - |
| <i>Bacillus cereus</i> | - | <i>Clostridium perfringens</i> | - | <i>Giardia intestinalis</i> WB Clone C6 | - | <i>Staphylococcus aureus</i> | - |
| <i>Bacteroides fragilis</i> | - | <i>Clostridium septicum</i> | - | <i>Giardia lamblia</i> | - | <i>Staphylococcus epidermidis</i> | - |
| <i>Campylobacter coli</i> | - | <i>Clostridium sordellii</i> | - | <i>Klebsiella oxytoca</i> | - | <i>Vibrio parahaemolyticus</i> | - |
| <i>Campylobacter fetus</i> subsp. <i>fetus</i> | - | <i>Clostridium sporogenes</i> | - | Norovirus | - | <i>Yersinia enterocolitica</i> | - |
| <i>Campylobacter jejuni</i> | - | <i>Cryptosporidium muris</i> | - | | - | | - |

14. Version history

| Version number | Chapter and designation |
|----------------|--|
| 2013-04-03 | Release version |
| 2018-10-19 | General revision |
| 2018-10-19 | 4. Reagents provided 5. Storage instructions 6. Additional necessary reagents and necessary equipment 9. Test procedure 10. Quality control 11. Result interpretation 13. Performance characteristics 14. Version history 15. Explanation of symbols |

15. Explanation of symbols

General symbols

| | |
|---|------------------------------|
|  | For in vitro diagnostic use |
|  | Consult instructions for use |
|  | Lot number |
|  | Expiry |
|  | Store at |
|  | Article number |
|  | Number of tests |
|  | Date of manufacture |
|  | Manufacturer |

Testspecific symbols

Not applicable

16. Literature

1. Kaper JM, *et al.* PATHOGENIC ESCHERICHIA COLI. Nature Reviews Microbiology 2004; 2:123-140.
2. Steffen R, *et al.* Vaccination against enterotoxigenic Escherichia coli, a cause of travelers' diarrhea. J Travel Med 2006; 12:102-107
3. Nataro JP, *et al.* Diarrheagenic Escherichia coli. Clinical Microbiology Reviews 1998; 11(1): 132-201.
4. Veira N, *et al.* High Prevalence of Enteroinvasive Escherichia Coli isolated in a remote region of Northern Coastal Ecuador. Am J Trop Med Hyg 2007, 76(3): 528-533.