

# ParoReal<sup>®</sup> Kit *Acanthamoeba* T4

Kit version 1.1

## Instructions for Use



CE

IVD

For *in vitro* diagnostic use

REF

DHUP00153

Σ

50 reactions



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## Explanation of symbols

|  |   |  |                                    |
|--|---|--|------------------------------------|
|  | Batch code  |  | Use by date                        |
|  | Catalogue number  |  | Manufacturer                       |
|  | Contains sufficient for <n> tests   |  | Temperature limit (Store at)       |
|  | This product fulfils the requirements of the European Directive 98/79 EC for <i>in vitro</i> diagnostic medical devices |  | In vitro diagnostic medical device |
|  | Consult instructions for use  |  | Unique device identifier           |
|  | Keep away from sunlight   |  | Contents                           |

## 1. Intended purpose

ParoReal® Kit *Acanthamoeba* T4 is a non-automated IVD real-time PCR test for the qualitative detection of DNA (18S rRNA gene) of *Acanthamoeba* species of genotype T4 (*A. castellani*, *A. lugdunensis*, *A. mauritaniensis*, *A. polyphaga*, *A. rhyodes*, *A. royreba*). T4 genotype is the most prevalent (approx. 86%) *Acanthamoeba* genotype causing keratitis worldwide. The test does not detect other *Acanthamoeba* genotypes (T3, T15, T11, and T5) also causing keratitis.

Proper specimens are DNA extracts isolated from human clinical specimens associated with keratitis (ocular swabs, corneal biopsies, ocular punctates, corneal scrapings) as well as contact lenses and contact lens solution.

This test is suitable for patients of all ages with suspected infection with *Acanthamoeba* genotype T4 (causative agent of *Acanthamoeba* keratitis, AK) and is intended as an aid in the diagnosis of infection with this pathogen in combination with patient history and additional clinical information.

The test is intended for professional use and is limited to qualified personnel instructed in the procedures of real-time PCR and *in vitro* diagnostic procedures.

## 2. Product description

ParoReal Kit *Acanthamoeba* T4 is a real-time PCR test and detects the 18S rRNA gene of *Acanthamoeba* species of genotype T4, which cause approx. 86% of *Acanthamoeba* keratitis worldwide (Diehl et al., 2021). The 18S rRNA gene of *Acanthamoeba* is a multicopy gene. The number of rRNA repeats in *Acanthamoeba* cells is 24 per haploid genome. However, because *Acanthamoeba* is polyploid, each cell contains approximately 600 rRNA genes (Yang et al., 1994).

A probe-specific amplification-curve in the FAM channel indicates the amplification of *Acanthamoeba* specific DNA. An internal DNA positive control (IPC) is detected in Cy5 and serves as a control for DNA extraction and possible real-time PCR inhibition. The target for the DNA IPC (artificial target DNA) is added during sample extraction.

This test has been validated with the ABI® 7500 Fast instrument (fast cycle parameters are not supported, Thermo Fisher Scientific) and was also tested with a LightCycler® 480 II (Roche Diagnostics), QuantStudio™ 7 real-time PCR system (Thermo Fisher Scientific), Mic instrument (bio molecular systems) and CFX Opus Real-time PCR System (Bio-Rad).

It is also compatible with other real-time PCR instruments which detect and differentiate fluorescence in FAM and Cy5 channel (e.g., QuantStudio™ 7 real-time PCR system (Thermo Fisher Scientific), qTOWER<sup>3</sup>G (Analytik Jena), cobas z 480 Analyzer (Roche)).

When using PCR-platforms not tested by ingenetix, an evaluation of the multiplex-PCR shall be done. Keep in mind that some PCR-platforms first must be calibrated with the corresponding dye before performing multiplex-PCR.

Ingenetix ParoReal®, BactoReal®, ViroReal®, MycoReal® and PanReal Kits are optimized to run under the same thermal cycling conditions. DNA and RNA can be analysed in one run.

## 3. Pathogen information

The genus *Acanthamoeba* is primarily a free-living protozoan in natural habitats, but also causative agent of human and animal disease. *Acanthamoeba* serves as host for a variety of pathogenic bacteria such as *Mycobacterium avium*. Infestation with *Acanthamoeba* is associated with potentially sight-threatening contact lens-related keratitis, serious infections of other organs and fatal granulomatous amoebic encephalitis. More than 20 species of *Acanthamoeba* are known, which can be classified into three morphologic groups (Group I, II and III) and 22 genotypes (T1-T22). Some species ((*A. castellanii* (T4), *A. lugdunensis* (T4), *A. polyphaga* (T4), *A. rhyodes* (T4), *A. quina* (T4), *A. palestinensis* (T2), *A. griffinii* (T3), *A. lenticulata* (T5), *A. astronyxis* (T7), *A. culbertsoni* (T10), *A. hatchetti* (T11), *A. healyii* (T12), *A. byersi* (T18), *A. divionensis*) have been recently associated with human disease. Genotype T4 has been considered the most important genotype in both ocular and CNS infections. According to literature, T4 genotype is the most prevalent *Acanthamoeba* genotype causing keratitis worldwide (approx. 86%) (Diehl et al., 2021).

## 4. Principle of real-time PCR

The test is based on multiplex real-time PCR by 5'-nuclease-assay technology. Specific DNA sequences are amplified and the generated PCR-products are detected by oligonucleotide-probes labelled with fluorescent dyes. This allows a sequence-specific detection of PCR amplicates.

During PCR, primers are extended by *Taq* polymerase and probes hybridized to the target are cleaved by the 5'-exonuclease activity of *Taq* polymerase. According to the accumulation of PCR product, the fluorescence of the probe increases with each PCR cycle. The change in fluorescence of the different dyes is recorded cycle by cycle in the real-time PCR instrument in the closed reaction tube at different fluorescence wavelengths.

The C<sub>q</sub> value (C<sub>q</sub> = Quantification cycle, C<sub>t</sub> = Cycle threshold, C<sub>p</sub> = Crossing point) describes the cycle at which the fluorescence first rises significantly above the background fluorescence.

## 5. Contents of the kit, stability, and storage

Table 1

| Labelling                                    | Content   | Amount      | Storage  |
|--|---|-------------|--|
| Acanthamoeba T4 + IPC3 Assay Mix (green cap) | Primers and probes for detection of<br>- <i>Acanthamoeba</i> (FAM)<br>- DNA IPC (Cy5) | 1 x 50 µl   | -25 to -15 °C  |
| IPC-Target (DNA) (orange cap)                | Target for DNA IPC (internal DNA positive control system)                             | 1 x 200 µl  | -25 to -15 °C  |
| Acanthamoeba T4 Positive Control (red cap)   | DNA positive control (approx. 1,000 target copies/µl)                                 | 1 x 300 µl  | -25 to -15 °C  |
| DNA Reaction Mix (white cap)                 | PCR reaction mix for DNA amplification  | 1 x 500 µl  | <b>-25 to -15 °C after first opening</b><br><b>2 to 8 °C</b> |
| Nuclease-free water (blue cap)               | Nuclease-free water   | 1 x 1000 µl | -25 to -15 °C  |

### DNA Reaction Mix

The Master Mix provided with the kit has been designed for reliable, high-sensitivity real-time PCR. The Master Mix contains a highly purified *Taq* Polymerase for rapid hot-start PCR, dNTPs with dUTP and Uracil-N glycosylase (UNG) to eliminate amplicon carryover, ROX™ dye (passive reference) and buffer components – additives optimized to handle RT-PCR inhibitors.

### Delivery and Storage

Shipment is with dry ice or with coolpacks. When stored properly, the kit components are stable until the specified expiration date. This also applies after opening. Store kit protected from light.

### Quality Control Release Testing

In accordance with the ISO 13485-certified Quality Management System of ingenetix, each lot is tested against predetermined specifications to ensure consistent product quality. Quality control is performed with a plasmid containing parts of the pathogen DNA.

## 6. Additionally required materials and devices

- Reagents and devices for DNA-extraction which are appropriate for the listed sample material (see 9. Preparation of the samples)
- Nuclease-free water
- Powder-free disposable gloves
- Pipettes (adjustable)
- Filter pipette tips
- Vortex-Mixer
- Benchtop centrifuge with rotor for 2 ml reaction tubes
- Real-time PCR instrument detecting and differentiating fluorescence in FAM and Cy5 channel
- Appropriate 96 well optical reaction plates or reaction tubes with corresponding (optical) closing material
- Optional: Laminar flow box
- Optional: PCR workstation

## 7. Precautions and safety information

### 7.1. General information

- For *in vitro* diagnostic use. The use of this kit is limited to qualified personnel instructed in real-time PCR and *in vitro* diagnostic procedures.
- Transportation of clinical specimens must comply with local regulations for the transport of biological materials.
- Specimens should be handled as if infectious in accordance with safe laboratory procedures. Wear protective powder-free disposable gloves when handling kit reagents and specimens.
- Discard specimens, reagents, and waste according to your local safety regulations.
- Improper collection, transport or storage of specimens may hinder the ability of the assay to detect the target sequences.
- Quality of DNA has a profound impact on the test performance. Ensure that the used DNA extraction system is compatible with real-time PCR technology.
- The real-time PCR instrument should be calibrated, serviced, and cleaned regularly.
- Protect kit components from light.
- Avoid mixing reagents of different kits and lots and check expiry date of the kit.

### 7.2. Specific information

A workflow must be followed to avoid false positive results due to detection of contaminating DNA.

#### Recommended measures to avoid DNA contamination:

- Use separated workspaces for specimen preparation, set-up of real-time PCR and amplification. Supplies and equipment must be dedicated to each of these separate workspaces to ensure workflow in the laboratory from pre- to post-PCR.
- Laboratory benches and devices must be cleaned regularly.
- Sample preparation should be performed in a laminar flow box. Clean laminar flow box regularly in all areas.
- If possible, leave consumables and pipettes in the laminar flow sterile bench and PCR workstation.
- Preparation of real-time PCR should be done in a PCR workstation.
- The use of sterile aerosol-resistant pipette tips is required.
- Use only DNA-free consumables.
- Wear lab coat.
- Work only with powder-free disposable gloves, do not touch the outer surface of the palm and fingers of the gloves when putting them on. Change gloves frequently. To avoid skin contact, wear gloves over the sleeves of the lab coat. Use disposable sleeve protectors if possible.
- Do not touch the rim or threads of open vials.
- Caution is advised when handling specimens and positive control to avoid cross-contamination.
- Store positive and potentially positive material separately from all other reagents.
- For a valid interpretation of results, a negative control shall be included during DNA-extraction (e.g., extraction of water instead of sample material), in to exclude false-positive results due to contamination with DNA of pathogen during extraction.
- Optional: include a negative control of PCR per PCR-run (nuclease-free water instead of sample, NTC).

## 8. Limitations

- Reliable results with this test are only achieved by appropriate specimen collection, transport, and storage, as well as an appropriate DNA extraction procedure.
- DNA extraction and *Acanthamoeba* detection have been validated for clinical ocular specimen material and contact lens material with this kit.
- A negative test result does not exclude the possibility of *Acanthamoeba* infection, because test results may be affected by improper specimen collection, technical error, specimen mix-up or pathogen quantities below the assay sensitivity. The presence of PCR inhibitors may lead to invalid results.
- For this kit highly specific primers and probes have been selected. However, false-negative, or less sensitive results might be obtained due to sequence heterogeneity within the target region of yet unknown clinical subtypes of genotype T4. The test detects *Acanthamoeba* species of genotype T4. Other *Acanthamoeba* genotypes (T3, T15, T11, and T5), which also cause keratitis, are not detected.
- Results should be interpreted in context of other clinical and laboratory findings.



## 9. Preparation of samples

ParoReal® Kit *Acanthamoeba* T4 is suitable for analysis of DNA extracts of samples from the human eye (ocular swabs, corneal biopsies, ocular punctates, corneal scrapings) as well as contact lenses and contact lens solution.

Sample preparation should be performed with the recommended measures to avoid DNA contamination (see precautions and safety information). Always include a DNA extraction negative control (e.g., extraction of water).

Purified DNA should be stored at -25 to -15 °C.

### 9.1. Sample collection and storage

Samples from the respiratory tract can be stored in microcentrifuge tubes.

Swabs can be collected with swab material proper for PCR (e.g., sterile polyester or rayon swabs with aluminium or plastic shaft) and put into 1 ml isotonic saline solution (NaCl 0.9%, not provided).

It is recommended to process samples immediately after collection. Store samples at 2-8 °C for no longer than 48 hours or freeze at -20/-80 °C.

### 9.2. Recommended extraction methods

Ensure that the extraction system used is not contaminated with DNA of pathogens detected with ParoReal® Kit *Acanthamoeba* T4. Extract the sample using a DNA extraction system compatible with real-time PCR technology and appropriate for the sample material.

#### For manual extraction recommended

- QIAamp DNA Mini Kit (Qiagen)
- High Pure PCR Template Preparation Kit (Roche)

When using extraction methods not recommended by ingenetix, an evaluation of the extraction method must be performed.

### 9.3. Quality control for DNA extraction and PCR inhibition with IPC

The DNA IPC system (internal DNA positive control) is used as a control for DNA extraction, identifies possible PCR inhibition and confirms the integrity of kit reagents.

For this, an artificial target DNA (IPC-Target (DNA), approx.  $6 \times 10^5$  copies/ $\mu$ l) is added during extraction.

**Note:** The Cq values of the IPC depend on the extraction method and on the type of sample material. Negative samples should show Cq values of the IPC between 27-30. The extraction method used must be validated accordingly with sample material. Use the IPC-Target (DNA) freshly diluted (1:10 with nuclease-free water) in the extraction, if Cq values < 27 are determined during validation with samples.

#### 9.3.1. Application of IPC as control of extraction and real-time PCR

The IPC-Target (DNA) must be added during extraction.

→ Per sample spike 1  $\mu$ l IPC-Target (DNA) (orange cap) directly into the appropriate volume of lysis buffer (or spike it to the sample after the lysis buffer has been added to the sample), then continue the extraction procedure.

**Caution:** The IPC-Target (DNA) shall not be added to sample material in the absence of lysis buffer, as degradation may occur. It must be added to the lysis buffer.

#### 9.3.2. Application of IPC as quality control for PCR reaction

If the IPC-Target (DNA) has not been added during extraction, it can be added at a later stage to the PCR master mix.

→ Freshly dilute the IPC-Target (DNA) 1:100 with nuclease-free water and add 1 µl of the dilution/PCR reaction (approx. 6000 target copies).

**Caution:** The IPC-Target (DNA) shall not be added to the master mix undiluted.

## 10. Preparation of real-time PCR

- Include one positive control (red cap), one extraction negative control and optionally one PCR negative control (NTC, e.g., nuclease-free water) per PCR run.
- It is generally recommended to analyse samples in duplicates, which increases the probability of pathogen detection and facilitates interpretation of results.
- Thaw DNA samples on ice.
- Thaw kit components completely at room temperature. When thawed, mix components carefully, centrifuge briefly with low speed.
- Mix the DNA Reaction Mix gently to ensure homogeneity of solution.
- **Positive Control**  
→ Use 9 µl of Positive Control (red cap). Always, pipette positive control at last.

### 10.1. Pipetting scheme

|  |                                  | <b>Per sample</b> |
|--|----------------------------------|-------------------|
| <b>Preparation of Master Mix</b><br>(mix well) | DNA Reaction Mix                 | 10.0 µl           |
|  | Acanthamoeba T4 + IPC3 Assay Mix | 1.0 µl            |
|  | <b>Total volume Master Mix</b>   | <b>11.0 µl</b>    |
| <b>Preparation of PCR</b>                      | Master Mix                       | 11.0 µl           |
|  | DNA-Sample*                      | 9.0 µl            |
|  | <b>Total volume</b>              | <b>20.0 µl</b>    |

\*1-9 µl of the sample can be used. For ≠ 9 µl sample, the volume must be adjusted with nuclease-free water.

→ **If IPC-Target (DNA) was not added during extraction:** Freshly dilute the IPC-Target (DNA) (orange cap) 1:100 with nuclease-free water and add 1 µl per sample directly to the master mix. In this case, the IPC is used for quality control of the PCR reaction. Only 8 µl of DNA sample can be analyzed.

**Caution:** The use of more than 1 µl diluted IPC-Target (DNA) per reaction might cause inhibition of the real-time PCR reaction.

- Prepare the Master Mix according to the number of samples, calculating an additional volume of approx. 10% for pipetting loss.
- Pipette 11 µl of the prepared Master Mix per sample into the well of the optical reaction plate.
- Then add 9 µl of the extracted sample or controls. Pipette the positive control at last.
- Seal the plate with a suitable optical sealing material.
- Vortex the sealed plate for 1-2 seconds and briefly centrifuge the plate.



## 10.2. Programming of temperature profile

Please find further information on programming of the real-time PCR instrument in the respective operator's manual. Keep in mind that some PCR-platforms must be calibrated with the corresponding dye before performing a multiplex-PCR.

**Sample Volume:** 20 µl

**Ramp speed:** "Standard" without "fast cycling" parameter for ABI® 7500 Fast Instrument, QuantStudio™ (Thermo Fisher Scientific)

| Program 1                   | Program 2                   | Program 3   |
|-----------------------------|-----------------------------|---|
| Cycles: 1<br>Analysis: None | Cycles: 1<br>Analysis: None | Cycles: 45<br>Analysis: Quantification<br>Acquisition at 60°C |
| 50 °C<br>2 min              | 95 °C<br>2 min              | 95 °C<br>5 sec<br>60 °C<br>1 min                              |

**Ad program 1:** If viral RNA should be also detected in the same PCR run, program 1 must be prolonged to 15 min at 50°C. This temperature profile can be used for all ingenetix ViroReal®, BactoReal®, MycoReal®, PanReal and ParoReal® kits for the detection of DNA or RNA.

**Ad program 2:** the previous temperature profile with 20 sec in program 2 can still be used.

### Select detection channels

**FAM channel:** Detection of *Acanthamoeba*

**Cy5 channel:** Detection of IPC

**For ABI® 7500 Instrument, QuantStudio™ 5/6/7 (Thermo Fisher Scientific), Mx3005P® (Agilent)**

FAM-TAMRA

Cy5-NONE

Passive reference dye: ROX

**For MIC Instrument (bio molecular systems)**

FAM Green

Cy5 Red

Passive reference dye: no ROX needed

**For cobas z 480 Analyzer / LightCycler® 480 II (Roche)**

Detection format: 2 Color Hydrolysis Probe

FAM: Excitation at 465 nm, Emission at 510 nm

Cy5: Excitation at 610 / 618 nm, Emission at 670 / 660 nm

Passive reference dye: no ROX needed

## 11. Interpretation of data

For the analysis of the PCR results, select the fluorescence display options FAM channel for the pathogen target and Cy5 channel for the IPC.

Please note that some PCR platforms require Color Compensation when using a multiplex PCR.

**IMPORTANT:** Samples should be checked in both logarithmic (Roche instrument: Abs Quant/Fit Points) and linear view and compared to the negative controls. In addition to the Cq values, also check the amplification curves and adjust the threshold (noise band) if necessary.

After you have saved the new settings, export the data. For the cobas z 480 Analyzer, export tables per dye.

Assessment of clinical specimen test results should be performed after the positive and negative controls have been examined and determined to be valid. If results of controls are not valid, no interpretation of results with clinical samples is possible.

### Negative samples:

Samples without amplification curves (no Cq values, undetermined) are regarded as negative. No DNA was detected in these samples because there is no infection with the corresponding pathogen, or the pathogen DNA concentration is below the detection limit of the test.

### Positive samples:

Samples with Cq values < 45 in the fluorescence channel for the pathogen target are regarded as positive.

**Table 2** shows the criteria for valid positive and negative controls.

Table 3 shows interpretation of data with clinical samples.

### 11.1. Controls

**Table 2** Criteria for valid controls, IPC-Target (DNA) was added during extraction

|                                    | Cq FAM channel<br><i>Acanthamoeba</i> target | Cq Cy5 channel<br>IPC <sup>1</sup> | Interpretation | Action   |
|------------------------------------|--|------------------------------------|----------------|----------|
| <b>Positive control</b>            | <29  | Negative                           | Valid          | -        |
| <b>Positive control</b>            | Negative                                     | Negative                           | Invalid        | See 12.1 |
| <b>Extraction negative control</b> | Negative                                     | 27-30 <sup>2</sup>                 | Valid          | -        |
| <b>Extraction negative control</b> | Negative                                     | Negative                           | Invalid        | See 12.1 |
| <b>Extraction negative control</b> | Positive                                     | 27-30 <sup>2</sup>                 | Invalid        | See 12.3 |
| <b>NTC PCR (optional)</b>          | Negative                                     | Negative                           | Valid          | -        |
| <b>NTC PCR (optional)</b>          | Positive                                     | Negative                           | Invalid        | See 12.2 |

<sup>1</sup> If the IPC-Target (DNA) was added directly to the master mix, all samples in the Cy5 channel must be positive.

<sup>2</sup> Cq values of the IPC are dependent on the extraction method and should show Cq values of 27-30 (see 9. preparation of samples).

## 11.2. Clinical samples

Samples with positive Cq-values are considered positive, see Table 3.

**Table 3** Interpretation of data with clinical samples

|                 | Cq FAM channel<br><i>Acanthamoeba</i> target | Cq Cy5 channel<br>IPC          | Interpretation | Action   |
|-----------------|--|--------------------------------|----------------|----------|
| Clinical sample | Negative                                     | 27-30 <sup>1</sup>             | Negative       | -        |
| Clinical sample | Positive                                     | Positive/Negative <sup>2</sup> | Positive       | -        |
| Clinical sample | Negative                                     | Negative                       | Invalid        | See 12.5 |

<sup>1</sup> A positive signal of the IPC excludes potential PCR inhibition. However, IPC Cq-values should show comparable results among samples. A shift of Cq- values can indicate a partial inhibition of PCR. Cq values of the IPC are dependent on the extraction method and should show Cq values of 27-30 (see 9. preparation of samples).

<sup>2</sup> High pathogen loads in the sample can lead to a reduced or absent fluorescence signal of the IPC.

In case of invalid data, analysis must be repeated with the remaining or newly extracted DNA sample (see 12. Troubleshooting).

## 12. Troubleshooting

### 12.1. No pathogen specific signal with positive control and with IPC

- Incorrect programming of the temperature profile or incorrect setting of detection channels on the real-time PCR instrument.
  - Compare temperature profile and setting of detection channels with details specified in the protocol
- Incorrect configuration of PCR reaction.
  - Check your pipetting steps with the pipetting scheme and repeat PCR, if necessary.
  - The DNA may have been degraded.
- The IPC-Target (DNA) was added undiluted directly to the master mix and not freshly diluted 1:100. The PCR reaction is therefore inhibited.
  - Freshly dilute IPC-Target (DNA) 1:100 and repeat PCR.
- No Positive Control was added.
  - Repeat PCR in case all clinical samples are negative.
- For control of real-time PCR only: 1 µl of freshly 1:100 diluted IPC-Target (DNA) must be added to the master mix. If the addition of IPC-Target has been forgotten:
  - Freshly dilute IPC-Target (DNA) and repeat PCR.
- For control of DNA extraction and PCR inhibition, the undiluted IPC-Target (DNA) must be added during extraction to the lysis buffer. If the addition of IPC-Target has been forgotten:
  - Repeat DNA extraction.

### 12.2. Pathogen signal with negative control

- If there are Cq values in the negative control below the cut-off specified in the Instructions for Use, a contamination occurred during the preparation of the PCR.
  - Repeat PCR with new reagents in replicates.
  - Strictly pipette positive control at last.
  - Make sure that workspace and instruments are cleaned at regular intervals.

### 12.3. Pathogen signal with extraction negative control

- If there are Cq values in the negative control of extraction below the cut-off specified in the Instructions for Use, a contamination occurred during the preparation of the PCR or the DNA extraction.
  - Repeat the PCR with unused reagents in replicates.
  - If repeatedly positive: Repeat DNA extraction and PCR using new reagents.
  - Strictly pipette positive control at last.
  - Make sure that workspace and instruments are cleaned at regular intervals and that workstations are separated from samples with high pathogen concentration.

**12.4. IPC specific signal with PCR negative control and positive control**

- The IPC-Target (DNA) has been added to lysis buffer during extraction, but there is IPC specific signal with PCR-NTC and positive control: Contamination with the IPC-Target (DNA).  
→ Make sure that workspace and instruments are cleaned at regular intervals.

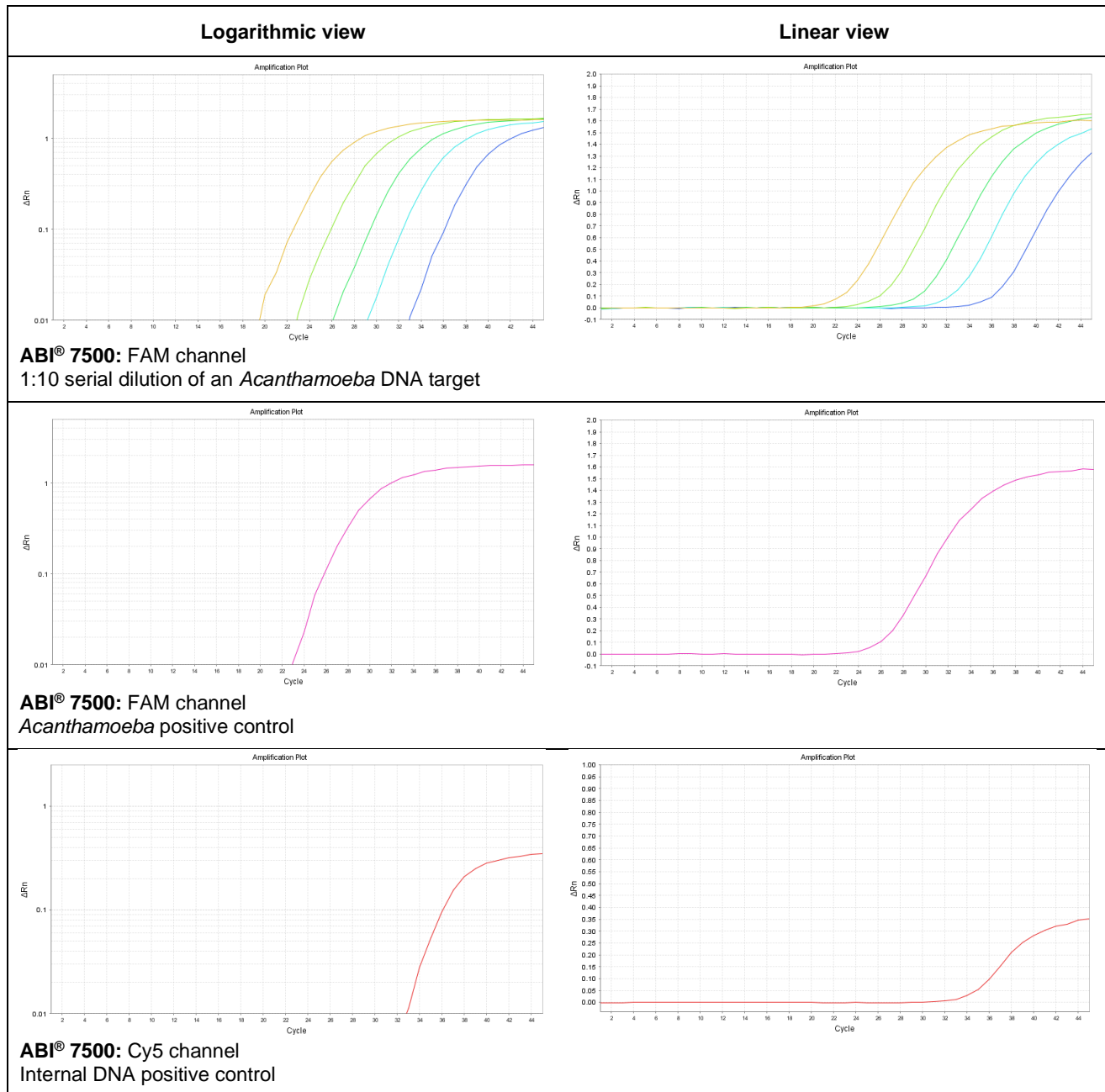
**12.5. No signal with IPC and no pathogen specific signal with sample**

- Incorrect assignment of detection channels in sample.  
→ Please verify the correct assignment of detection channels.
- The DNA might be degraded.
- If the IPC-Target (DNA) was added during extraction:
  - Inhibition of PCR may have occurred.
  - DNA extraction was unsuccessful.
  - The IPC-Target (DNA) was not added to the lysis buffer of the sample.
  - The extracted sample was not added to the PCR-reaction.  
→ No statement is possible. Verify you use a recommended method for DNA isolation and re-examine the single steps of the DNA extraction.  
→ If no operating mistakes during DNA extractions can be retraced, it is recommended to repeat the PCR with lower amounts of DNA-eluate (1/5 or 1/10 of sample volume + the adequate volume of nuclease-free water).

### 13. Specifications and performance evaluation

#### 13.1. Test performance on different real-time PCR instruments

Performance of ParoReal® Kit *Acanthamoeba* T4 with an ABI® 7500 instrument (Thermo Fisher Scientific) is shown in Figure 1.



**Figure 1** Performance of ParoReal® Kit *Acanthamoeba* T4

This test has been validated with the ABI® 7500 instrument (Thermo Fisher Scientific) and was also tested with a LightCycler® 480 Instrument II (Roche), QuantStudio™ 7 real-time PCR system (Thermo Fisher Scientific), Mic instrument (bio molecular systems) and CFX Opus Real-time PCR System (Bio-Rad).

### 13.2. Limit of detection, LoD95%

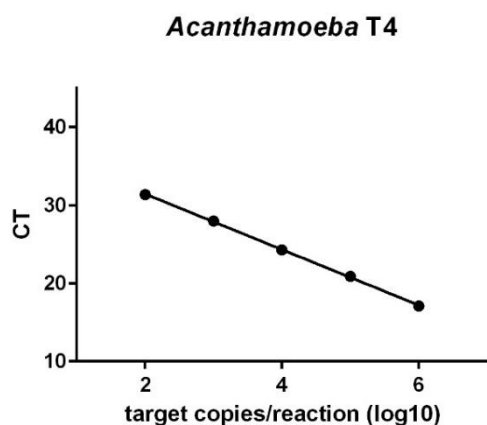
**Method:** The limit of detection (LoD95% = smallest number of copies of target DNA which can be detected in 95% of cases) was determined by testing a commercially available CE certified reference material with known concentrations of *A. castellanii* (AMPLIRUN® *Acanthamoeba castellanii* DNA control (Vircell, order No MBC054). Twenty replicates of at four different concentrations around the detection limit were tested (0.1, 0.5, 1, 3 copies). Calculation was performed with a non-linear (logistic) curve fit using GraphPad Prism Software.

**Result:** The LoD95% is 0.5 genome copies/reaction. The 18S rRNA gene is a multicopy gene and is present up to 22 times in the haploid genome of *Acanthamoeba*.

### 13.3. Linearity and dynamic measuring range

**Method:** ParoReal® Kit *Acanthamoeba* T4 was tested with a 10-fold dilution series of a plasmid containing a fragment of *Acanthamoeba* DNA. Linearity was determined using a 10-fold dilution series ( $10^6$  –  $10^1$  target copies/reaction) of the plasmid. The number of determinations (n) per dilution was nine.

**Result:** The assay shows linearity over the range of 100 to 1,000,000 target copies/reaction with a slope of  $3.561 \pm 0.04463$  and an  $R^2$  of  $> 0.99$  as shown in Figure 2.



**Figure 2** Ten-fold dilution series of an *Acanthamoeba* DNA standard

### 13.4. Precision

**Method:** Precision within a run (intra-assay), between multiple runs (inter-assay) and between two lots (inter-lot) was determined.

**Result:** The mean values of the coefficients of variation (CV%) are 1.2% for intra-assay precision, 1.2% for inter-assay precision and 0.5% for inter-lot precision.

### 13.5. Analytical specificity

**Method BLAST analysis:** The selection of highly specific primers and probes ensures analytical specificity. The specificity of primer and probes was validated *in silico* using the Basic Local Alignment Search Tool (BLAST) against the NCBI database. Primers and probes have been checked for possible homologies to currently published sequences. This analysis validates the detection of so far known strains of *Acanthamoeba* species of genotype T4.

**Result:** This test detects the 18S rDNA gene of *Acanthamoeba* species of genotype T4 (*A. castellanii*, *A. lugdunensis*, *A. mauritaniensis*, *A. polyphaga*, *A. rhyodes*, *A. royreba*). No relevant sequence variabilities were observed in the primer and probe region thereof. The test does not detect other *Acanthamoeba* genotypes (T3, T15, T11, and T5) also causing keratitis, due to mismatches in primer and/or the probe. There is no cross-reactivity to other organisms.



### 13.6. Diagnostic evaluation

#### Method:

Diagnostic evaluation of ParoReal® Kit *Acanthamoeba* T4 was performed by an external service with 168 DNA isolates from clinical specimens. Samples were DNA isolates from ocular swabs (n=63), corneal biopsies (n=20), ocular punctates (n=7), corneal scrapings (n=43) as well as contact lenses (n=7) and contact lens solution (n=26) and other material (n=2). They represented 67 samples positively tested for *Acanthamoeba* and 101 *Acanthamoeba*-negative samples. DNA was isolated from clinical specimens using the Qiagen QiaAmp Mini Kit (n=29) or Roche High Pure PCR Template Preparation Kit (n=139).

Results were compared with those obtained with a DIN EN ISO 15189 accredited pathogen-specific quantitative real-time PCR reference method detecting the 18S rRNA gene of the genus *Acanthamoeba* using hybridization probes. Real-time PCR was performed using a LightCycler® 480 II (Roche Diagnostics).

#### Result:

Out of the 67 samples which were positive for *Acanthamoeba* with the reference method, 59 samples were also positive with ParoReal® Kit *Acanthamoeba* T4 (true positive). Cq values were between 19-37. Six out of the remaining 8 samples which were positive with the reference method but negative with ParoReal® Kit *Acanthamoeba* T4 were sequenced and phylogenetically identified by BLAST analyses in the NCBI database. These samples were identified as *Acanthamoeba* T3 genotype. The 101 DNA samples which were negative with the reference method were also negative with ParoReal® Kit *Acanthamoeba* T4 (true negative).

The diagnostic evaluation of ParoReal® Kit *Acanthamoeba* T4 is shown in Table 4 and Table 5.

**Table 4** Overall results obtained with 168 tested clinical samples, 2x2 contingency table

|                                      | Reference |     |       | Total |
|--------------------------------------|-----------|-----|-------|-------|
|                                      | pos       | neg | Total |       |
| ParoReal® Kit <i>Acanthamoeba</i> T4 | pos       | 59  | 0     | 59    |
|                                      | neg       | 8*  | 101   | 109   |
| Total                                | 67        | 101 | 168   |       |

\* Six samples were identified as *Acanthamoeba* T3 genotype.

**Table 5** Statistical evaluation of the diagnostic validation

|             | Value   | 95% CI            |
|-------------|---------|-------------------|
| Sensitivity | 88.06%  | 77.82% to 94.70%  |
| Specificity | 100.00% | 96.41% to 100.00% |
| NPV         | 92.66%  | 86.82% to 96.03%  |
| PPV         | 100.00% |                   |
| Prevalence  | 39.88%  | 32.42% to 47.71%  |

## 14. References

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- Lorenzo-Morales, Jacob; Khan, Naveed A.; Walochnik, Julia. 2015. An update on *Acanthamoeba* keratitis: diagnosis, pathogenesis and treatment. *Parasite* 22: 10.
- Yang, Q., Zwick, M. G., & Paule, M. R. (1994). Sequence organization of the *Acanthamoeba* rRNA intergenic spacer: identification of transcriptional enhancers. *Nucleic acids research*, 22(22), 4798–4805.

## 15. Revision history

| Revision | Date        | Description  |
|----------|-------------|--|
| 1.0      | 25 May 2022 | First version  |
| 1.1      | 01 May 2023 | <p><b>Change of the company address</b></p> <p><b>5. Contents, stability and storage</b><br/>Kit version 1.1:<br/>Filling volume of IPC-Target (DNA) and Positive Control was changed.<br/>The DNA IPC Target is not stored in RNA/DNA Stabilizer anymore and was renamed to IPC-Target (DNA).</p> <p><b>7. Precautions and safety instructions</b><br/>Update of this section<br/>Inclusion of additional information on specimen collection and storage, processing of specimens after collection, storage conditions of specimens, transport and storage conditions of purified DNA.<br/>Recommended measures to avoid DNA contamination added.</p> <p><b>9. Preparation of samples</b><br/>Update of paragraph of Quality control for RNA extraction and PCR inhibition<br/>Note that for Cq values &lt;27, the IPC-Target (DNA) must be diluted prior to extraction.</p> <p><b>11. Interpretation of PCR data</b><br/>Table 1 and Table 2: Update</p> |

### Note:

Any serious incident that has occurred in relation to the product shall be reported to the manufacturer and the competent authority of the Member State in which the user and/or the patient is established.

### Technical support

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