



## Instructions For Use

# LightMix<sup>®</sup> Modular Enterovirus

500

Cat.-No. 50-0656-96

Roche SAP n° 07 730 454 001

Kit with reagents for 96 PCR reactions 20 µl for detection of Enterovirus [lyophilized]

## 1. Content Storage and Expiry

- 1 Vial orange cap 96 reactions Enterovirus (lyophilized)
- 1 Vial black cap Positive Control (RNA), Cp-value ~ 31

## Storage at Arrival:

Store cooled or at ambient temperature  
Do **not** freeze the lyophilized reagents.

- Lyophilized kits are stable for at least 6 months (4°C to 25°C in the dark). See lot-specific expiry date.
- Dissolved reagents are stable for at least 2 weeks if stored protected from light and cooled (4°C).
- Dissolved reagents can be stored long-term at -20°C (within expiry). Avoid multiple freeze-thaw cycles.
- Dissolved positive controls must be stored at -20°C. Avoid multiple freeze-thaw cycles.

## 2. Additional Reagents required

LightCycler<sup>®</sup> Multiplex RNA Virus Master

Cat.-No. 06 754 155 001

## 3. Introduction

Enteroviruses (EV) have a positive-sense ssRNA genome and belong to the Picorna viruses, which are classified based on pathogenesis into Poliovirus (causing poliomyelitis), Coxsackie A, B, and Echovirus.

EV infections are common and are associated with several diseases, in particular respiratory infections ('summer cold') and gastrointestinal disease. EV is spread mainly through the fecal-oral route.

This kit detects the human Enterovirus species A, B (including Echovirus), C (including Poliovirus) and D (including EV-D68); Coxsackieviruses are members of species A, B and C. Other members of the Picornaviridae, in particular Rhinovirus (HRV) and Parechovirus (hPeV) are not detected. For specific detection of EV-D68 we offer the ModularDx kit 53-0122-96.

## 4. Description

A 80 bp fragment from the 5-UTR region is amplified with specific primers and detected with a Cyan500 labeled hydrolysis probe.

## 5. Specification

This assay detects 10 genome equivalent copies or less per reaction (in vitro transcribed RNA).

## 6. Sample Material and Extraction

Typical specimen are from feces or rectal swabs (gastrointestinal infections) or nasopharyngeal swabs (respiratory disease). See ModularDx Document **Extraction Protocols**.

## 7. Material Safety Data (MSDS)

According to OSHA 29CFR1910.1200, Australia [NOHSC:1005, 1008 (1999)] and the EU Directives 67/548/EC and 1999/45/EC any products which do not contain more than 1% of a component classified as hazardous or classified as carcinogenic do not require a Material Safety Data Sheet (MSDS).

Product is not hazardous, not toxic, not IATA-restricted. Product is not from human, animal or plant origin. Product contains synthetic oligonucleotide primers and probes.



## 8. Instructions for Use

- Instrument programming see document
- Color Compensation see instructions in
- Pipetting instructions multiplex PCR see

**ModularDx Programming**

**40-0320 Universal Color Compensation Hexaplex**

**ModularDx Multiplex**

### 8.1. Programming Roche 480 Instruments

See the Instrument operator's manual for details. Start programming before preparing the solutions. The protocol consists of four program steps:

- 1: Reverse Transcription of the viral RNA
- 2: Denaturation: sample denaturation and enzyme activation
- 3: Cycling: PCR-amplification
- 4: Cooling: cooling the instrument

#### Detection Format 500 Channel

LightCycler® 480 Instrument:

LightCycler® 480 II Instrument:

cobas z 480 Analyzer (open channel):

**Set Quant Factor 10, Max Integration time 1 sec**

450-500

440-488

**No filter combination for Cyan500 (opt. use FAM channel)**

| Program Step:               | RT Step     | Denaturation | Cycling             |               |          | Cooling  |
|-----------------------------|-------------|--------------|---------------------|---------------|----------|----------|
| <b>Parameter</b>            |             |              |                     |               |          |          |
| Analysis Mode               | <b>None</b> | None         | Quantification mode |               |          | None     |
| Cycles                      | 1           | 1            | 45                  |               |          | 1        |
| Target [°C]                 | 55          | 95           | 95                  | 60            | 72       | 40       |
| Hold [hh:mm:ss]             | 00:05:00    | 00:05:00     | 00:00:05            | 00:00:15      | 00:00:15 | 00:00:30 |
| Ramp Rate [°C/s] <b>96</b>  | 4.4         | 4.4          | 4.4                 | 2.2           | 4.4      | 1.5      |
| Ramp Rate [°C/s] <b>384</b> | 4.6         | 4.6          | 4.6                 | 2.4           | 4.6      | 2.0      |
| Acquisition Mode            | None        | None         | None                | <b>Single</b> | None     | None     |

Table 1

### 8.2. Experimental Protocol

- **Sample material:** Use aqueous nucleic acid preparations (e.g. 'High Pure Viral Nucleic Acid Kit').
- **Negative control:** Always run at least one no-template control (NTC) - replace the template DNA with water.
- **Positive control:** Run a positive control - replace the template DNA with the provided Positive Control.

For an increased sensitivity use 10 µl sample per 20 µl reaction, in case that inhibition is likely to occur, e.g. extracts obtained from fecal samples, use 5 µl. For 10 µl reactions in 384 well plates use 5 µl /2.5 µl.

#### 8.2.1. Preparation of Parameter-Specific Reagents (PSR, 96 reactions):

One reagent vial with a **orange** cap contains all primers and probe to run 96+ LightCycler® reactions.

**Add 50 µl** PCR-grade water to each reagent vial, mix the solution (vortex) and spin down.

For robotic pipetting the volume can be extended to 55 µl (signals will decrease by 10-20%).

► **Use 0.5 µl** reagent for a 20 µl PCR reaction.

#### 8.2.2. Preparation of the Positive Control

**Add 160 µl** RNase/DNase-free Tris buffer or water to the vial with the **black** cap, for 10 µl sample add **320 µl**. Mix the solution by pipetting up and down 10 times. If vortexing spin down to collect the solution.

**Notes:** Opening of this vial may cause contaminations of the work-space (aerosol). Use of Tris buffer pH 8.0-8.5 increases the long-term stability in solution. Store dissolved controls frozen.

► **Use 5 µl** positive control (≈ 1,000 copies) for a 20 µl PCR reaction (or 10 µl if using 10 µl sample).

### 8.2.3. Preparation of the Reaction Mix

Multiply volumes by the number of reactions plus controls and one reserve and prepare in a cooled tube:

| For use with the Roche LightCycler® Multiplex RNA Virus Master |  |                |
|--|--|----------------|
| for 5 µl extract   | Component  | 10 µl extract  |
| 10.4 µl  | <b>Water</b> , PCR-grade (colorless cap, provided with the Roche Master kit)   | 5.4 µl         |
| 0.5 µl   | <b>Reagent mix</b> (parameter specific reagents containing primers and probes) | 0.5 µl         |
| --   | Control Reaction and additional assays (Multiplex PCR)                         | --             |
| 4.0 µl   | <b>Roche Master</b> (see Roche manual)   | 4.0 µl         |
| 0.1 µl   | <b>RT Enzyme</b> (see Roche manual)  | 0.1 µl         |
| <b>15.0 µl</b>   | <b>Volume of Reaction Mix</b>  | <b>10.0 µl</b> |

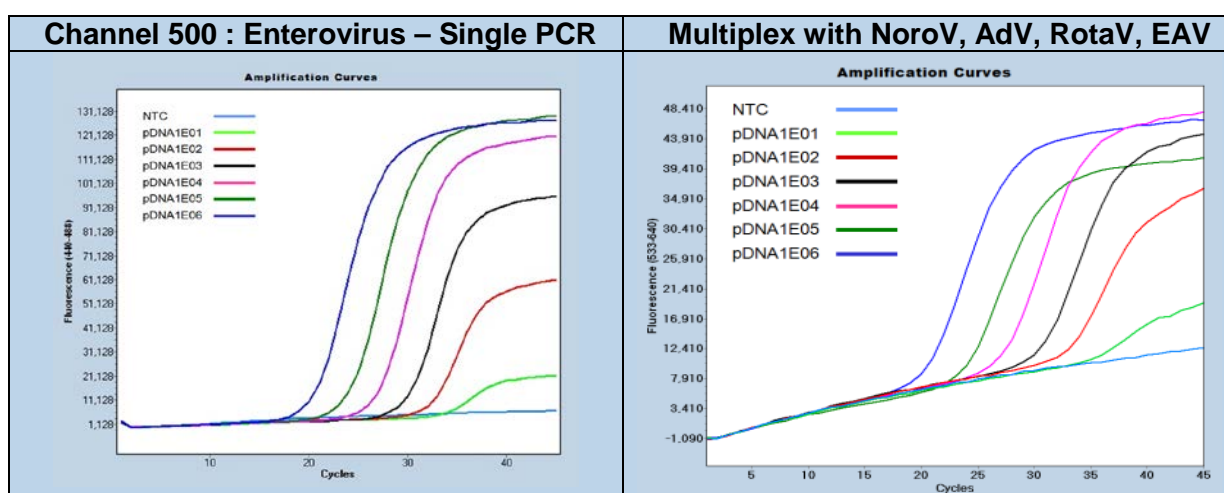
Table 2

Mix gently, spin down and **transfer 15 µl (10 µl)** per well.

**Add 5 µ (10 µl)** of sample or control to each well for a final reaction volume of 20 µl. Seal plate and centrifuge.

**Start run**

### 9. Typical Results (Data from LightCycler® 480 II system)



Dilution row 1E6 to 10 copies / reaction

Figure 1

### 10. Reading the Results

Perform data analysis as described in the operator's manual. For multiplex assays select the color compensation. We recommend using the Second Derivative Maximum method (Automated (F" max). View results in the 500 channel. The negative control (NTC) must show no signal.

| Channel 500 (sample)   | Channel 660 Control Reaction | Channel 500 NTC Control | Result               |
|------------------------|------------------------------|-------------------------|----------------------|
| No amplification       | Detectable                   | Negative                | Not detectable       |
| Amplification Cp < 37+ | Not relevant                 | Negative                | Enterovirus Positive |
| No amplification       | Not detectable               | Not relevant            | PCR failure Repeat   |
| Amplification signal   | Not relevant                 | Positive                | Contamination Repeat |

**Notes:** cobas z 480 Analyzer signal levels are about 50% compared to LightCycler® 480 II results.

+ Recommendation : Define the cut-off 2-4 cycles higher than observed for 10 copies.

### 11. References

The development, implementation and evaluation of a Real-Time PCR-based diagnostic service for viral causes of infectious intestinal disease. Gunson, 2007

## 12. Multiplex PCR Compatibility Respiratory Virus and Gastro Virus Panel

This Enterovirus assay can be combined with other assays up to 6plex reactions including an internal control (IC) or a spiked extraction control (for example EAV) as depicted below :

**Multiplex PCR and Instrument Compatibility**  
Color Compensation 40-0320 is mandatory for Multiplex PCR


| 500      | 530     | 580     | 610     | 640        | 660       |
|----------|---------|---------|---------|------------|-----------|
| EnterovV |         |         |         |            |           |
| EnterovV | InfA    | InfB    | RSV-A/B | HRV        | MSTN or   |
| EnterovV | PeV     | MPV     | AdV     | HRV        | PhHV or   |
| EnterovV | GG1+GG2 | RotaV A | AdV     | Astrovirus | EAV or    |
| EnterovV | GG1+GG2 | RotaV A | AdV F   | Astrovirus | Roche RPC |

| 480 II | z 480 | LC96 | LC2.0 | Nano |
|--------|-------|------|-------|------|
| X      | X     | X    |       |      |
| X      |       |      |       |      |
| X      |       |      |       |      |
| X      |       |      |       |      |
| X      |       |      |       |      |

Table 3

## 13. Version History

|         |  |            |
|---------|--|------------|
| V140909 | Release version  | 2014-09-09 |
| V151001 | 2015 protocol Multiplex Master, 5/10 µl extract and 60°C acquisition | 2015-10-01 |
| V160313 | 1. Storage of controls, 8.2.2 buffer, 8.2.3 wording                  | 2016-05-21 |

| <b>Certificate of Analysis (CoA)</b>  |        |     |     |          |     |     |  |
|---|--------|-----|-----|----------|-----|-----|---|
| Lot n°<br>Expiry :  |        |     |     |          |     |     |   |
| <b>Dilution</b>   | 1E6    | 1E5 | 1E4 | PC       | 1E2 | 1E1 | <b>passed</b>   |
| <b>Cp range</b>   |        |     |     |          |     |     |   |
| <b>Measured</b>   | 29-31  |     |     |          |     |     |   |
| <b>Signal level</b>   |        |     |     |          |     |     |   |
| <b>Measured</b>   | 80-120 |     |     |          |     |     |   |
| <b>Negatives</b>  | 10/10  |     |     |          |     |     |   |
| <b>Note:</b> Cp (crossing point) values collected with pDNA (single target PCR). Fluorescence (FL) levels depend on instrument settings and may vary. The Cp values will vary from instrument to instrument by up to 2 cycles, while the distance between two dilution steps should be relative constant (ΔCp). |        |     |     |          |     |     |   |
| <b>QC Acceptance Date:</b>  |        |     |     | YYYYMMDD |     |     |   |
| We, the undersigned, certify that the product designated above has been obtained in accordance with the rules of production and quality control.  |        |     |     |          |     |     |   |
| <b>Name(s) :</b>  |        |     |     |          |     |     |   |

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