#### DESCRIPTION

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**Evolving** 

**Enzymes** 

Innovating IVD

TOROIVD<sup>®</sup> Probe 1-step RT-qPCR 5G Premix is a fast, single-tube, 2×RT-qPCR mix that provides for sensitive, reproducible detection up to four RNA/DNA targets in a single multiplex reaction. Particularly useful for virus detection with TaqMan<sup>®</sup> probe assays, the mix includes thermostable MMLV reverse transcriptase, TOROIVD<sup>®</sup> 5G DNA polymerase, dNTPs and reaction buffer all in a single tube. The improved enzymes and reaction mixture combination also enables a high resistance to PCR inhibitors and high stability in room temperature. The 1-step system is suitable for high-throughput analysis because of its simple reaction setup. In addition, this system can reduce the risk of cross-contamination. The premix is suitable for high-speed RT-qPCR and enables accurate detection and quantification of targets, making it possible to obtain highly reproducible and reliable real-time PCR results over a wide dynamic range.

#### **FEATURES**

#### -Rapid and highly sensitive

This kit can achieve the rapid and highly sensitive quantification of a low-copy targets by a one-step RT-qPCR method with probes and be suitable for the quantification of RNA/DNA viruses or mRNA expressed at a low level.

#### -Optimized for multiplexing

This kit has been validated for multiplexing up to five targets simultaneously, allowing for additional targets and/or controls to be run simultaneously for efficiency or quality control purposes.

#### -Inhibitor tolerant

The unique proprietary formulation of this kit allows robust performance even in the presence of substances that can normally inhibit PCR, such as heparin, hematin, or EDTA, increasing your confidence when working with a variety of complex clinical samples.

#### -Wide dynamic range compatible with RNA and DNA

This kit has been optimized to provide high specificity and dynamic range for use with both RNA and DNA targets. This input flexibility can help streamline the number of different workflows in your lab to improve efficiency.

#### -Broad instrument compatibility

This kit can be run in either fast or standard cycling conditions with equivalent performance across a wide variety of real-time cyclers. The  $50 \times ROX$  Reference dye (not supplied) is added and can be applied to the real-time cyclers that require a passive reference dye.

#### -Utilization of dUTP

This kit contains dUTP in the reaction buffer . Therefore, the rate of false-positive detection can be reduced by adding UNG (not supplied).

#### **COMPONENTS**

The kit includes the following reagents, and QPR-300 can be used for 200 reactions for a total  $25\mu$ l reaction volume. All reagents should be stored at  $-20^{\circ}$ C.

Cat NO.	Components	Size
QPR-300	2× RT-qPCR 5G Premix	1.25mL×2 tubes/bag

#### Notes:

 $-2 \times \text{RT-qPCR}$  5G Premix contains TOROIVD<sup>®</sup>III reverse transcriptase, RNase inhibitor and TOROIVD<sup>®</sup>5G DNA polymerase, 0.4mM dA/C/G/T/UTP, 5mM Mg<sup>2+</sup>, reaction buffer and stabilizer, etc.

#### **NOT SUPPLIED**

In some experimental applications, the following reagents may be used with QPR-300, which are not supplied in this kit. Please contact us to order.

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# **TOROIVD®** Probe 1-step RT-qPCR 5G Premix

Cat NO.	Components	Size
UNG002YL	Heat-labile Uracil-N-glycoslyase	200µl/tube; 1ml/tube; 10ml/tube.
ROX-050	50×ROX Reference dye	100µl/tube, 1ml/tube, 10ml/tube.
RDB-100	RNA Dilution&Storage Solution	100ml/pcs,10L/pcs

#### Notes:

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-The  $50 \times ROX$  reference dyes are used for analyses with instruments that correct for cross-talk between wells, such as the real- time PCR instruments by Applied Biosystems and Agilent Technologies.  $0.5\mu$ l  $50 \times ROX$  Reference Dye was added for a total  $25\mu$ l reaction volume in when using the following instruments, Applied Biosystems 7300/7700/7900HT, StepOnePlus, etc. And  $0.05 \mu$ l was added for using the following instruments, Applied Biosystems 7500/7500Fast Step OnePlus ,Agilent Technologies AriaMx, etc. No ROX Reference Dye is required when using other brand instruments, such as LightCycler 96/LightCycler 480 system (Roche), CFX96 Real-Time PCR Detection System (Bio-Rad), Smart Cycler System (Cepheid) ,etc. -RNA Dilution&Storage Solution is a buffer that provides greater RNA stability than TE Buffer, RNase-free water and RNA Virus VTM medium. RNA Dilution&Storage Solution is compatible with direct 1- step RT-qPCR.

#### **PRIMER/PROBE DESIGN**

#### -Design of primers

Primer length: 18–25bp; Tm of primer: 60–65°C; GC content: 40–60%; Purification grade: HPLC grade; Target length: 70–200 bp; Larger targets (>200 bp) tend to reduce the efficiency and specificity of amplification. **-Design of probes** 

Probe length: 20–30bp; Tm of probe: 65–70°C; GC content: 40–60%; Purification grade: HPLC.

#### -Checking the performance of primers and probes:

-Prepare a dilution series with five or more dilutions of template RNA/DNA. Perform RT-qPCR assay using the diluted RNA/DNA with the newly designed primers and probe, and draw a standard curve.

- Confirm that the PCR efficiency is between 90% and 110% and  $R^2$  is equal to or greater than 0.99. if the PCR efficiency or  $R^2$  are outside of these ranges, the reaction conditions should be optimized. If this does not improve the result, the primers and/or probe should be redesigned.

#### PROTOCOL

1. This kit should be fully thawed before use. Gently vortexed and briefly centrifuged.

- 2. Purified or crude template RNA/DNA can be may be used directly or after dilution.
- 3. Prepare the following reaction mixture in a thin-walled qPCR tube or plate.

Components	25μL reaction	
2× RT-qPCR 5G Premix	12.5µL	
Heat-labile Uracil-N-glycoslyase	0.1-0.3µL	
10µM Forward primer	1µL	Premix
10µM Reverse primer	1µL	Treninx
10µM TaqMan® probe	0.4µL	
50×ROX	0/0.05/0.5µL	
DNase/Rnase Free Water	XuL	
Template RNA/DNA solution	5µL	

4. Gently mix the reaction solutions and spin down in microcentrifuge.

#### Notes:

- 2.5mM MgCl2 in final concentration have been added in this reaction mixture. But for the direct RT-qPCR to crude template RNA, the MgCl<sub>2</sub> concentration may need to be optimized between 2.5-8mM of final concentration.

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BSA and Triton X-100 may need to be added to improve the performance of direct RT-qPCR. -The primer concentration should be optimized between 0.2-0.8  $\mu$ M and TaqMan<sup>®</sup> probe optimized between 0.1-0.4  $\mu$ M with 10-50 copies templates /reaction. So the best primers-probe concentration sets was selected by orthogonal design of experiments.

-UNG should be used and not supplied with this kit. The added volume is according to the instructions of the manufacturers.

# **CYCLING CONDITIONS**

The recommended 2-step PCR protocol is described below:

For ABI 7500/7300 etc.				
Steps		Temperature	Time	Cycles
1	Reverse transcription	52°C	5min	1
2	Prenaturation	95℃	1min	1
3	Denaturation	95°C	10 sec	40-45
	Annealing/ Extension	60°C	30 sec	

For Bio-Rad CFX96,ABI StepOne Plus,etc.				
	Steps	Temperature	Time	Cycles
1	Reverse transcription	52°C	5min	1
2	Prenaturation	95℃	1min	1
3	Denaturation	98°C	3 sec	40-45
	Annealing/ Extension	60°C	10 sec	

For Bioer LineGene 9600 Plus, Roche LightCycler 96 /LightCycler 480 systems,etc.				
Steps		Temperature	Time	Cycles
1	Reverse transcription	52°C	5min	1
2	Prenaturation	95℃	1min	1
3	Denaturation	95°C	10 sec	40-45
	Annealing/ Extension	60°C	20 sec	

# Notes:

-Use this protocol first and optimize PCR conditions when necessary. Perform 3-step PCR when using primers with low Tm values or when 2-step PCR is not feasible.

-The indicated UNG treatment temperature can be optimized 25-37°C, and time between 0-5min.

-The indicated RT temperature can be optimized between 50-60°C and time between 2-15min.

-The indicated Pre-denaturation temperature can be optimized 95-98°C, and time between 2min-5min.

-The indicated denaturation temperature can be optimized 95-98°C, and time between 3sec-10sec.

-The indicated Extension /Annealing temperature can be optimized  $60-65^{\circ}C$ , and time between 5sec-30sec. Fluorescence signal gathering should be set up at this step.

# **APPLICATION DATA**

# Template: MS2RNA from Roche

# **Primer and Probe:**

Forward primer:GCCTTAGCAGTGCCCTGTCT 400nM Reverse primer:AACATGCTCGAGGGCCTTA 400nM Taqman Probe:FAM-CCCGTGGGATGCTCCTACATGTCA-TAMRA 200nM Reagents:

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TOROIVD Probe 1-step RT-qPCR 5G Premix (QPR300)

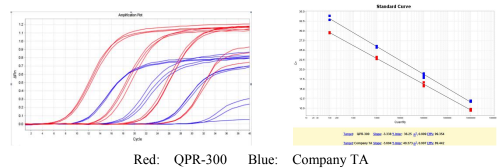
# Example 1. Comparison of the dynamic range and PCR efficiency with company TA MS2RNA dilutions:

1.58×10<sup>2</sup>;1.58×10<sup>4</sup>;1.58×10<sup>6</sup>;1.58×10<sup>8</sup>;

#### Instrument:

ABI 7500

**Results:** 



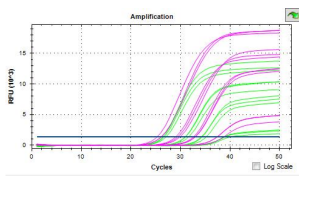
Example 2. Sensitivity comparison with company TB. MS2RNA dilutions:

 $1.58 \times 10^{0}; 1.58 \times 10^{1}; 1.58 \times 10^{2}; 1.58 \times 10^{3};$ 

# Instrument:

CFX 96

**Results:** 



Pink: QPR-300 Green: Company TB

# **STORAGE**

This reagent can be stored at 4°C for 2 months. For longer storage, this reagent should be kept at -20°C for 2 years $_{\circ}$