ABScript II One Step RT-qPCR Probe Kit



Catalog: RK20407

Size: 20 RXN / 100 RXN (50 µL/RXN)

2X One Step RT-qPCR Probe	RM21462
Buffer	
One Step Probe HS Taq	RM21463
ABScript II RT Enzyme Mix	RM21464
50X ROX Dye I	RM21465
50X ROX Dye II	RM21466
Nuclease-free H ₂ O	RM20214

Product Description

ABScript II One Step RT-qPCR Probe Kit a ready-to-use kit allowing reverse transcription and subsequent probe-based qPCR in a single tube. It contains all components for RT-qPCR except primers, probes and RNA templates. The one-step format significantly improves sensitivity and effectively prevent contamination. The ABScript II Reverse Transcriptase in the kit provides reliable reverse transcription to a wide range of RNA template amount. After reverse transcription, the Hot-start version of Tag polymerase is activated at 95°C and the ABScript II Reverse Transcriptase is inactivated simultaneously. In the sequential PCR reaction, the 5 '-3 'exonuclease activity of Tag polymerase cleaves the hybridized probe, separating the reporter from the quencher and releasing fluorescent signal. The ABScript II One Step RT-qPCR Probe Kit is an ideal product for high-speed analyses of low input RNA sample.

Product Components

Component	20 RXN (50 μL / RXN)	100 RXN (50 μL / RXN)
2X One Step RT-qPCR	500 µL	1.25 mL X 2
Probe Buffer *		
One Step Probe HS Taq	20 µL	100 µL
ABScript II RT Enzyme Mix**	20 µL	100 µL
50X ROX Dye I ***	20 µL	100 μL
50X ROX Dye II ***	20 µL	100 μL
RNase-free ddH₂O	500 µL	1.25 mL X 2

- * Containing dNTPs, Mg²⁺, etc.
- ** Containing ABScript II Reverse Transcriptase, RNase Inhibitor
- *** Passive reference dye to normalize the fluorescence signals

Storage

Upon receipt, store all components at -20°C.

Compatible Instruments

50X ROX Reference Dye I

Applied Biosystems 7000/7300/7700/7900, Applied Biosystems StepOne[™]/StepOnePlus[™]

50X ROX Reference Dye II

Applied Biosystems 7500/ViiA7TM, QuantStudio[™], Stratagene Real-time PCR Systems, Rotor-gene[™] 3000

NO ROX Reference Dye

Bio-Rad iCyclers/ CFX96/ CFX 384, Roche Light Cyclers[®], QIAGEN/Corbett Systems, Eppendor Mastercyclers[®]

Additional Material

Required but not Supplied

- 1. RNA templates, primers and probes
- Optical-grade qPCR tubes, plates, sealing films, and aerosol-resistant pipette tips

Precautions

- Fully thaw the 2X One Step RT-qPCR Probe Buffer before use. Mix the buffer well and avoid directly sunlight.
- Determine the total number of reactions required and prepare master mix. Triple replicates for each reaction are recommended.
- 3. The ABScript II RT Enzyme Mix and One Step Probe HS *Taq* contain high concentration of glycerin. Mix gently before use without generating air bubbles. Spin briefly to collect all the contents at the bottom. After use, return it to -20°C immediately.
- 4. If applicable, use aerosol-resistant pipette tips and microtubes to minimize contamination.
- High quality RNA templates are recommended for optimal results.
- Only gene specific primers are recommended.
 Random primers and Oligo dT primers are NOT recommended in the reverse transcription reaction
- 7. The optimal length of amplicon is between 70 and 200 bp for general cycling condition.

Protocol

Prepare materials before reaction setup:

- Pipette, aerosol-resistant pipette tip, cold blocks and ice.
- Gene expression primers and probes
- RNA templates
- 1.5 mL RNase–free EP tubes, Real–time PCR tubes and plates

Set up One Step RT-qPCR

experiment

1.Prepare the reaction mix

Set up the reaction on ice by adding the following components for the number of reactions required.

Component	Volume	Volume
2X One Step RT-qPCR	10 μL	25 µL
Probe Buffer		
One Step Probe HS Taq	0.4 µL	1 µL
ABScript II RT Enzyme Mix	0.4 µL	1 µL
Forward Primer (10 µM) *	0.4 µLeach	1 µL each
Reverse Primer (10 μM) *	0.4 µLeach	1 µL each
TaqMan Probe (10 μM) ***	0.4 µLeach	1µL each
50X ROX Dye (As require by	0.4 µL	1 μL
instrument guideline)		

Total RNA **	Up to 2 μL	Up to 5 µL
RNase-free H₂O	Up to 20 μL	Up to 50 µL

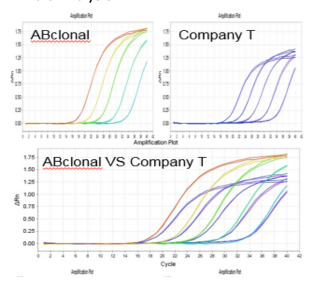
^{*} A final primer concentration of 0.2 μ M is recommended for most reactions. However, to optimize individual reaction, a primer titration from 0.1 μ M to 1.0 μ M can be performed. The length of amplified PCR products should ideally be in the range of 70 – 200 bp.

Optimized One Step RT-qPCR program:

Step	Temperature	Time	Cycles
Reverse Transcription	42℃	5 min	1
Polymerase Activation	95℃	3 min	1
Denaturation,	95℃	5–15 s	
Annealing and Extension	60°C	30–34 s*	40

^{*}The extension time should be adjusted to the minimum time required for data acquisition according to qPCR instrument guidelines used. (30 s for Applied Biosystems StepOnePlus™, 31 s for Applied Biosystems 7300, and 34 s for Applied Biosystems 7500)

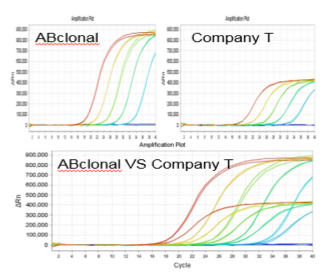
2.Data Analysis



Pic 1: Amplification plots comparing ABScript II One Step RT-qPCR Probe Kit and one-step RT-qPCR kit from Company T. Rat total RNA in ten-fold dilution from 150 ng to 15 pg was used as template. Identical RT-qPCR programs were run with Applied Biosystems StepOnePlus[™] to detect VIC fluorogenic probe. The ABclonal kit outperformed the kit from competitor T with higher detection sensitivity.

^{**} Use 15 pg~150 ng of RNA template in a 20 µL reaction.

^{***} A Probe concentration of 50-250 nM is recommended.



Pic 2: Amplification plots comparing ABScript II One Step RT-qPCR Probe Kit and one-step RT-qPCR kit from Company T. Rat total RNA in ten-fold dilution from 150 ng to 15 pg was used as template. Identical RT-qPCR programs were run with Applied Biosystems StepOnePlusTM to detect FAM fluorogenic probe. The ABclonal kit outperformed the kit from competitor T with higher detection sensitivity.