

SuScript 1-Step RT PCR Kit

Catalog No Size • RT01A035 50 Rxn 200 Rxn • RT01A036

Description:

SuScript 1-Step RT PCR Kit is a convenient one-step solution for the RT-PCR reactions containing all the components necessary to obtain cDNA in a high yield from RNA samples and PCR amplification kit with hotstart activity. The RNase Inhibitor presented in the kit adequately protects RNA templates against degradation. Optimized reaction buffer, MgCl₂, Oligo-dTs, random hexamers and reverse transcriptase are also included in the 5X RTMix. The first strand of cDNA can be directly used as a template in downstream applications (PCR, Real-Time PCR, etc).

Kit Components:

Components	RT01A035	RT01A036
5X RT Mix	200 µl	800 µl
2X HS SuPCR Mastermix	1 ml	4 ml

General Considerations:

- High purity template RNA is essential for reliable efficient cDNA synthesis. A A260 / A280 ratio of 1.7 or higher is strongly recommended.
- The amount of template RNA is depended on the expected copy number of the sequence of interest. In general, 1 μg - 1 ng of total RNA is recommended, 0.05 - 100 ng if you are working with isolated mRNA.
- This protocol recommends cDNA synthesis for 1 hour at 42°C.
- To enhance the template coverage, the 5X RTMix also contains random hexamer primers. This provides multiple priming sites along the RNA for the detection of multiple short sequences.

Recommended Protocol:

This protocol is exemplary for one reaction, and for multiple reactions it is necessary to calculate the components in a proportional manner. Mix the kit components in a micro tube in the recommended proportions below. Briefly centrifuge the tube to spin down the contents and to eliminate any air bubbles. Place the tube on ice until to reverse transcriptase reaction.

Reaction Components	Volume	
5X RT Mix	4 µl	
Total RNA	1-6 µl	
DNase/RNase Free Water	Up to 20 µl	
Total Volume	20 µl	

Mix gently and carry-out reverse transcriptase reaction conditions:

RT Steps	Temp (°C)	Time	Cycle
RT Reaction	42	1 hour	1
RT Deactivation	80	10 min	1

Notice: The synthesized cDNA can be used immediately, without purification, or stored at -20 °C for future use.

Reaction Setup:

- It is recommended that thaw the 2X HS SuPCR Mastermix and mix thoroughly but gently to ensure even distribution of the components.
- Put the specimen's DNA on the PCR tubes or multi-well
- For negative control sample do not add DNA please use nuclease-free water.
- Keep plate or PCR tubes on ice during reaction set up.

Components	Volume	
2X HS SuPCR Mastermix	25 µl	
Forward primer (10 µM)	1 µl	
Reverse primer (10 µM)	1 µl	
Sample DNA (max 10 µl)	2 µl	
Nuclease-free water	X to complete 50 μl	
TOTAL Volume	50 μl	

PCR Conditions:

Steps	Temp.	Time	Cycles
HotStart Activation	95 ℃	5 min	1
	95 ℃	30 sec	
Amplification	50-60 ℃	30 sec	30-35
	72 ℃	30 sec (1 min / kb)	30-33
Final extension	72 ℃	2 min	1

Storage Conditions:

Store all contents at -20 °C in a freezer.

Quality Control:

Nicking activity, priming activity, exonuclease activity, or endonuclease activity has not been detected.

Evaluations of Results:

Analyze the amplification of the samples in gel electrophoresis, positive and negative controls for any band observed. If the amplicon band is observed in Positive Control but doesn't in Negative Control, amplification reaction was completed properly. If any other situation occurs, however, amplification reaction may be proceeding in a wrong way. In such a case, re-test the samples from reagent preparation or check the primer design.



SuGenomik Biyoteknoloji Ltd Şti

Address: METU Technopolis, Ankara.

Web: www.sugenomik.com

Mail: info@sugenomik.com

Tel: +90(312)385 85 89



Web: www.sugenomik.com

Mail: info@sugenomik.com

Tel: +90(312)385 85 89





