[Cat. No.] Please refer to the Ordering Information

Introduction

AccuPower[®] RocketScript[™] RT-PCR PreMix, RNase H Minus is a one-step RT-PCR product that uses *RocketScript[™]* Reverse Transcriptase (RTase), RNase H Minus and *ProFi Taq* DNA Polymerase independently developed by BIONEER. *RocketScript[™]* RTase, RNase H Minus is a recombinant *M-MLV* RTase with enhanced thermostability up to 70°C, as well as eliminated RNase H activity. It can produce efficiently full-length cDNA from long RNA transcripts up to 12.5 kb and synthesize cDNA even with small amounts of 1 pg of total RNA. In addition, included *ProFi Taq* DNA Polymerase provides accurate amplification and higher fidelity for PCR. This product contains vacuum-dried components including *RocketScript[™]* RTase, reaction buffer, DTT, dNTPs, RNase inhibitor, and *ProFi Taq* DNA Polymerase. It simplifies preparation of reverse transcription reaction mixture and PCR mixture by adding template RNA and primers without any extra process.

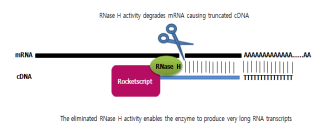




Figure 1. Elimination of RNase H activity. Eliminated RNase H activity from RTase complex enables the RTase to produce longer RNA transcripts, up to 12.5 kb, compared with original *RocketScript*™ RTase itself.

Applications

- Standard RT-PCR
- Long kb RT-PCR
- Virus detection
- Gene expression analysis

Features & Benefits

- Thermostable activity: *RocketScript*[™] RTase has excellent thermal resistance even at 70°C, so it is suitable for cDNA synthesis of secondary RNA structures.
- No RNase H activity: Useful for cDNA synthesis up to 12.5 kb from long target RNA using enzymes developed by proprietary genetic engineering technology.
- User-friendly: Simple amplification of long-size RNA difficult to be done with conventional one-step RT-PCR methods.

Composition

Composition	Concentration	
<i>RocketScript</i> ™ Reverse Transcriptase, RNase H Minus	200 U	
5X Reaction buffer	1X	
DTT	0.25 mM	
dNTPs (dATP, dCTP, dGTP, dTTP)	Each 250 µM	
RNase inhibitor	1 U	
ProFi Taq DNA Polymerase	1 U	
Stabilizer and tracking dye	1X	

Specifications

RocketScript™ Reverse Transcriptase, RNase H Minus			
DNase activity	No		
RNase activity	No		
RNase H activity	No		
Fragment size	Up to 12.5 kb		

Storage

Store at -20°C. If stored in the recommended temperature, this product will be stable until the expiration date printed out on the label.

Online Resources



Korean



English

Visit our **product page** for additional information and protocols

Ordering Information

Description			Cat. No.
0.2 ml thin-wall 96 tubes 8-tube strips with attached cap 480 tubes	20 µl/rxn	K-2231	
	90 lubes	50 µl/rxn	K-2233
	480 tubes	20 µl/rxn	K-2232
		50 µl/rxn	K-2234

Notice

BIONEER corporation reserves the right to make corrections, modifications, improvements and other changes to its products, services, specifications or product descriptions at any time without notice.

Explanation of Symbols



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Experimental Procedures

	Steps		Procedure Details			
	Preparation of reaction mixture	 Add template RNA, primers and nuclease-free water into AccuPower[®] RocketScript[™] RT-PCR PreMix, RNase H Minus tubes to make a total volume of 20 µl or 50 µl. Do not include the dried pellet. Amount of template RNA and primers 				
1		Comp	Components		20 µl reaction 50 µl reaction	
		Template RNA	Total RNA Poly(A) RNA	0.01-5 µg 0.01-5 µg	0.01-5 µg 0.01-5 µg	
		Primers	Gene specific primer	10-30 pmol	10-50 pmol	
		 Dissolve the vacuum-dried pellet by pipetting or vortexing, and briefly spin down. 3. Perform the reaction under the following conditions. 				
2	RT-PCR	Step	Temperature	Time	Cycles	
		cDNA synthesis	42-70°C	30 min	1 cycle	
		Pre-denaturation	95°C	5 min	1 cycle	
		Denaturation	95°C	10-30 sec		
		Annealing	55°C	10-30 sec	30-35 cycles	
		Extension	72°C	1 kb/min		
		Final extension	72°C	5 min	1 cycle	
		* Note: Reaction temperature should be optimized according to Tm value of primers.				
3	Analyze with gel electrophoresis	 4. After the reaction, maintain the reaction mixture at 4°C. The samples can be stored at -20°C until use. 5. Load samples on agarose gel without adding a loading-dye mixture, and perform gel electrophoresis for analysis. 				

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