

4× EZscript Reverse Transcription Mix II (with gDNA Remover)

Catalog No.: EZB-RT2GQ

Description

The **EZBioscience® 4× EZscript Reverse Transcription Mix II (with gDNA Remover)** is a new-generation reverse transcription kit with higher reverse transcription efficiency compared to previous generations.

The kit is suitable for real-time RT-PCR (RT-qPCR) that contains a gDNA Remover which can effectively eliminate the contamination of genomic DNA (gDNA) or other double stranded DNA during the analysis of gene expression. To accurately analyze gene expression, it is necessary to detect cDNA in samples without contaminating DNA. To avoid amplification of gDNA, primers can be designed on different exons spanning introns. However, there may be cases where a suitable primer cannot be designed, as with a gene with a single exon or a gene without a long intron. Also, it may be difficult to avoid unexpected amplification from gDNA due to non-specific amplification or the existence of pseudo-genes. Moreover, some labs are heavily contaminated by the PCR products of previous tests. This kit offers a potent gDNA Remover that can eliminate double stranded DNA in RNA sample in 5 minutes at room temperature without loss of RNA. Then the first strand of cDNA is synthesized by adding the 4× EZscript RT Mix II. Reaction products are applicable to subsequent PCR, qPCR.

The 4× EZscript RT Mix II contains reverse transcriptase, RNase Inhibitor, optimized buffer system, dNTPs, and Oligo dT18 and Random Hexamer as primers. The reverse transcriptase in this Mix is a genetic engineered enzyme based on M-MLV (RNase H-) reverse transcriptase. And the multiple site-mutations of reverse transcriptase can obviously increase its affinity to RNA templates and its strand extending ability, which make reverse transcription reaction more efficient. Moreover, this transcriptase is rather resistant to common reverse transcriptase inhibitors. At the same time, the kit uses the latest optimized reaction system to further improve the reverse transcription efficiency. This product is also very suitable for reverse transcription using plant RNA.

Components

Components	EZB-RT2GQ (100 Rxns)	EZB-RT2GQ-L (500 Rxns)
4× EZscript RT Mix II	550 µl	550 µl × 5 tubes

gDNA Remover	220 µl	220 µl × 5 tubes
Nuclease free ddH ₂ O	1 ml	1 ml × 5 tubes

Storage

Store at -20°C.

Caution

Avoid RNase contamination

Please keep the environment of experiment clean. Clean gloves and mask should be worn during the experiment. Centrifuge tubes, tips and other supplies used in the experiment must be RNase-free.

Protocol

gDNA Remover treatment of RNA

1. Add 1 µg total RNA (10 pg ~ 2 µg adjustable) or 200 ng Poly(A)⁺ RNA (10 pg ~ 500 ng adjustable) to a new RNase-free centrifuge tube. Add 2 µl gDNA Remover to the RNA, pipette up and down for 10 times to mix thoroughly. Incubate at room temperature (19 ~ 27°C) for 5 minutes.

Reverse Transcription

2. Add components to the gDNA Remover-treated RNA according to the following table, then mix gently with a pipette and centrifuge the mixture briefly to the bottom of the tube.

Components	20 µl Reaction
gDNA Remover treated RNA	X µl
4× EZscript RT Mix II	5 µl
Nuclease free ddH ₂ O	up to 20 µl

3. Perform the reverse transcription reaction at 42°C for 15 minutes and 95°C for 30 seconds.

4. The reverse transcription reaction product can be directly used in PCR applications, but the cDNA is recommended to dilute and mix thoroughly before use (The specific dilution factor depends on the abundance of gene expression. Generally, cDNA is diluted 5 ~ 10 times). If the qPCR experiment is not performed immediately, it is recommended to store at -80°C for long-term storage. Avoid repeated freeze-thaw cycles.