Enzymax LLC

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Product Information

Product Name: RNA Ligation/decapping enzyme Mix Order: info@enzymax.net Catalog #: LDM50 Order: info@enzymax.net Tel: 859-219-8482

Size: 500U (50 μ l) Fax: 859-219-0653 Web: www.enzymax.net

Product: RNA Ligation/decapping enzyme Mix

Product Description:

The RNA ligation and decapping enzyme mix is specially formulated and optimized for once-step RNA ligation and decapping in RNA ligase-mediated (RLM) 5' RACE (Rapid Amplification of cDNA Ends) and RNA library cloning or preparation for sequencing. The mixture of both enzymes enables the decapping and ligation reaction in a single step which simplifies the procedure. Below is the description for enzyme usage in decapping/ligation step from our RNA RACE kit (KIT96).

Components:

60 μl 10x ligation and decapping buffer	
120 μl 50% PEG-8000	
60 μl 10 mM MnCl ₂	
50 μl ligase and decapping enzyme mix	

Enzyme usage & RNA RACE procedure:

RNase inhibitor may be needed if samples have RNase contamination. Use of RNA inhibitor is recommended.

Step 1: 5' decapping and ligation

Step 1. 5 decapping and figation	
~1 µg total RNA*	3.5 µl
RNA linker	1 μl
Enzyme mixture	1 μl
PEG-8000 (mix well in the reaction)	2.5 µl
MnCl ₂	1 μl
10x buffer	1 μl

Assemble the above components in a DNase/RNase free tube and incubate the tube at room temperature for 2-8 h.

Step 2: Reverse transcription (RT)

Step 2. Ite i cise transering	otion (Iti)
Reverse primer (10 μM) **	0.5 µl
dNTP	1 μl
H ₂ O	4 ul

Add above components into step 1; incubate at 65 °C for 15 min, then keep on ice for 2 min.

Add components below and incubate at 42 °C for 30 min.

RT dilution buffer	2 μ1
DTT	1 μl
Reverse transcriptase	0.5 µl

^{**}This is a gene specific reverse primer, which is not included in the kit. However, it can be substituted by random hexamer. When using random hexamers (use 20 µM instead), incubate

samples at room temperature for 10 min before RT. Contact us for **global 5'RACE kit**.

Step 3: PCR and nested PCR:

NOTE: Except for the common primer, all other PCR reagents are not included in this kit. PCR reagents are sold separately. The following procedures are based on our tests and need to be optimized by customers.

1. PCR Reaction (1st PCR reaction): 50 μl

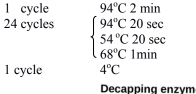
	A00
10xPCR buffer	5 µl
25 mM dNTP	$0.6 \mu l$
10 μM 5' common primer (outer primer)	1 μl
10 μM 3'gene specific primer (outer prime	r) 0.5 µl
RT product from step 2	1 μl
PCR polymerase	0.5 μl
H ₂ O	41.4 µl
1 avala 0.4° C 2 min	

l cycle	94°C 2 min
15 cycles	94°C 20 sec 54°C 20 sec 68°C 1min
	54 °C 20 sec
	68°C 1min
1 cycle	$4^{\circ}C$

2. Nested PCR Reaction: 50 µl

2. Trested I est Redection. 30 µs	
10x PCR buffer	5 μl
25 mM dNTP	0.6 µl
10 μM 5' common primer (outer primer)	1 μl
10 μM 3' nested gene specific primer **	* 1 µl
PCR product from 1 st PCR reaction	1 μl
PCR polymerase	0.5 µl
H_2O	40.9 μl

***Also called inner primer which is designed at upstream of 3'gene specific primer (outer primer). Instead of provided 5'common primer, you can also design inner forward primer, which is at downstream of 5'common primer. Link for primer design: http://www.clcsupport.com/clcgenomicsworkbench/650/ Nested PCR.html.



Decapping enzyme
- + ++

1000 bp

500 bp

5' RACE product

5' RACE product form different amount of decapping enzyme

^{*} RNA isolated by column, phenol, or Trizol-based methods are suitable for this kit. If RNA is partially degraded, it may be necessary to pre-treat sample with phosphatase (Calf Intestinal Phosphatase or CIP). RNA samples should be EDTA-free.