

Phi-Script™ 1st Strand cDNA Synthesis Kit

#### **Product Overview**

Phi (\$) Script <sup>™</sup> 1st strand cDNA synthesis contains PhiScript reverse transcriptase manufactured by dx/dt. It's an engineered form of Moloney Murine Lukemia Virus reverse transcriptase (M-MuLV-RT). Point mutations are introduced in \$\$Cript<sup>™</sup> reverse transcriptase so as to reduce RNAse H Activity and to improve its thermal stability. Good cDNA yields can be obtained when reverse transcribed at 42-55°C.

### **Catalog Details**

R6201 50 Reactions R6202 200 Reactions

### Pack Contains

1. Extrapure dNTPs (10μM Each)
2. Random Primers (50μM)
3. Oligo(dT) (50μM)
4. RNase Inhibitor (40U/μL)

### Store all @ -20°C

5. PhiScript RT (200U/µL)

- 6.5X RT Reaction Buffer
- 7. DTT (100mM)
- 8. Nuclease Free Water

### First Strand cDNA Synthesis Protocol

The most important element is - good quality RNA. Make sure you have extracted quality RNA (Total RNA or mRNA) from appropriate cells . The quality of the RNA can be checked using agarose gel electrophoresis to see two distinct rRNA. It is important to have intact RNA specially when longer, full length cDNA is expected. Make sure to have all the tips, plastic wares, other consumables and working area RNAse Free.

		Full length cDNA synthesis		Gene Specific cDNA Synthesis	
Components		20µL Read	ction Target	20µL Reaction	Target
Total RNA Template		variable	≤ 2 µg	variable	≤ 2 µg
dNTP Mix (10mM Each)		lμL	0.5 mM Each	lμL	0.5 mM Each
Either OR OR	Random Primers (50µM Stock Oligo(dT) primers (50µM Each Both 1µL Each	) ) 1µL	upto 200 ng - Random primers upto 500 ng - (oligo-dT)	0	NA
Gene Specific Primer(10µM Stock) (Reverse primer ONLY)		0	NA	lμL	2 - 20pmoles
Total Volume		ΧµL		X µL	

## PRE - RT STEP

Mix all the above componens (1. RNA, 2. dNTP, 3. Random primers OR oligo(dT) or both ) and heat it up at 70°C for 5 minutes. This step will remove secondary structures of RNA. After this, keep the vial on ice immediately at least for 1 minute. For gene specific assays, pre-RT step is optional, may not be essential.



# Phi-Script™ 1st Strand cDNA Synthesis Kit

### **RT STEP**

Perform the following reaction on ice.

Components	20µL reaction	Target Final Concentration
Pre - RT mixture	ΧμL	
DX/DT Rnase Inhibitor (40U/µL)	lμL	10 - 40U
5X φScript™ Reaction Buffer	4 µL	١Χ
DTT (100mM Stock)	lμL	Upto 2 - 10mM
φScript™ Reverse Transcriptase (200U/μL)	lμL	upto 3 µL
Nuclease Free Water	upto 20µL	

• Incubate the above mixture at 42°C for 30 - 60 minutes. Increase the reaction mixture temperature to 50-55°C for gene specific assays. If you are using random primers, then first incubate 25°C for 10 minutes prior to 42°C.

- Inactivate RT by heating at 85 ℃ for 10 minutes.
- Store the cDNA samples at -20°C for further analysis. This cDNA now can be used as PCR template for amplification.

### PCR STEP

We recommend to use DX/DT Extremo Mastermix (R2020/ R2021) for below PCR assays.

Components	20µL reaction	50µL reaction	Final Concentration	
Template DNA/cDNA	lμL	2 µL	<b>5ng</b> - 100 ng	
Forward Primer (10µM)	0.5 µL	lμL	0.1 - 1µM	
Reverse Primer ( 10µM)	0.5 µL	lμL	0.1 - 1µM	
Extremo MasterMix (2X)	10 µL	25 µL	1X	
Nuclease Free Water	Upto 20 µL	Upto 50 µL		

#### PCR Conditions

Step	Temperature	Time	Cycle
Initial denaturation	95 <b>℃</b>	2 minutes	1
Denaturation	95 <b>℃</b>	20-30s	
Annealing (Tm of your primers)	°C	20-30s	25 - 45 Cycles
Extension	72 ℃	1 minute/kb	
Final Extension	72 ℃	2 - 5 minutes	1
Hold, if required.	2 - 8 °C	variable	683

# Any Help / Contact



info@dxbidt.com |. +91-7349708807 | www.dxbidt.com

Scan the QR Code to read on trouble-shooting guide