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UltraScript™ cDNA Synthesis Kit



Product description:

The UltraScript™ cDNA Synthesis Kit uses the latest developments in reverse transcriptase technology and buffer chemistry to enhance cDNA synthesis speed and yield with accurate transcript representation. The reverse transcriptase, buffer system and combination of random hexamers with anchored oligo(dT) allow for unbiased, efficient and sensitive cDNA synthesis.

The modified MMLV reverse transcriptase (RTase) is both thermostable and extremely active. The RTase is not inhibited by ribosomal and transfer RNAs making total RNA an ideal substrate. The enzyme is blended with RNase inhibitor preventing degradation of RNA by contaminating RNase.

5x buffer contains anchored oligo(dT), random hexamers, enhancers, dNTPs and ${\rm MgCl_2}$. The relative concentrations of random hexamers and anchored oligo(dT) have been optimised for the generation of cDNA for use in real-time PCR experiments. The kit can be used with 4.0 pg to 0.4 μ g total RNA or oligo(dT) purified mRNA.

Component	25 reactions	100 reactions
5x cDNA Synthesis Mix	1 x 100 μL	1 x 400 μL
20x UltraScript™ for cDNA Synthesis	1 x 25 μL	1 x 100 μL

Shipping and storage

On arrival the kit should be stored between -30 °C and -15 °C. If stored correctly the kit will retain full activity for 12 months. Avoid exposure of the stock solution to frequent temperature changes and limit handling at room temperature to the necessary minimum. Do not store the mix once it is combined with the RTase.

Limitations of product use

The product may be used for in vitro research purposes only.

Technical support

Help and support is available on our website at https://pcrbio.com/resources/ including answers to frequently asked technical questions. For technical support and troubleshooting you can submit a technical enquiry online, or alternatively email technical@pcrbio.com with the following information:

- Reaction setup
- PCR cycling conditions
- Screen grabs of gel images / real-time PCR traces

Important considerations

5x cDNA Synthesis Mix: Contains anchored oligo(dT), random hexamers, 15 mM MgCl $_2$, 5 mM dNTPs, enhancers and stabilizers. It is not recommended to add further enhancers or MgCl $_2$ to the reaction. The buffer composition has been optimised to generate cDNA for downstream real-time PCR analysis.

Template: Use 4.0 pg to $0.4 \text{ }\mu\text{g}$ total RNA or oligo(dT) purified mRNA. For template amounts greater than $0.4 \text{ }\mu\text{g}$ we recommend our UltraScript 2.0 cDNA Synthesis Kits.

Incubation temperature: We recommend incubating with a temperature of 42 °C for 30 minutes for the majority of applications (<65% GC). Where regions of interest contain high secondary structure (>65% GC) incubation temperatures of up to 55 °C may be used.

qPCR setup: Users can add the cDNA created directly to a qPCR reaction, or dilute it 10x - 50x in PCR grade H_2O to reduce the concentration and extend the volume. We recommend adding 2.0 - 4.0 μL of cDNA solution to a 20 μL qPCR reaction.

Reaction setup

- 1. Allow 5x cDNA Synthesis Mix to thaw, briefly vortex.
- 2. Prepare a master mix based on the following table. Insert reagents in sequence listed:

Reagent	20 μL reaction	Final concentration	Notes
5x cDNA Synthesis Mix	4.0 µL	1x	
20x UltraScript™ for cDNA Synthesis	1.0 μL		Add before total RNA as RNase inhibitor is blended with RTase
Total RNA or oligo(dT) purified mRNA (between 4.0 pg and 0.4 μ g)	X μL		
PCR grade dH ₂ O	Up to 20 µL final volu	me	

No RT control setup (optional)

3. Prepare a master mix based on the following table. Insert reagents in sequence listed:

Reagent	20 µL reaction	Final concentration	Notes
5x cDNA Synthesis Mix	4.0 μL	1x	
Total RNA or oligo(dT) purified mRNA (between 4.0 pg and 0.4 µg)	Χ μL		Use equal amount as in step 2
PCR grade dH ₂ O	Up to 20 μL final volume		

Incubation and enzyme denaturation

- 4. Incubate at 42 °C for 30 minutes
- 5. Incubate at 85 °C for 10 minutes to denature RTase