# **RT-qPCR Extraction Control**

	Catalog numbers		
Batch : See vial	MDX028-1	500 Rxn	RT-qPCR Extraction Control Red
	MDX029-1	500 Rxn	RT-qPCR Extraction Control Orange

#### Storage and stability:

RT-qPCR Extraction Control is shipped on dry/blue ice. All kit components should be stored at -80 °C upon receipt. Excessive freeze/thawing is not recommended.

#### Expiry

When stored under the recommended conditions and handled correctly, quality is retained until the expiry date on the outer box label.

#### Quality Control:

The RT-qPCR Extraction Control is extensively tested for quality and the absence of contamination.

#### Safety Precautions:

Please refer to the material safety data sheet for further information.

#### Notes

This reagent has been manufactured under 13485 Quality Management System, and is suitable for research use only.

Monitoring of RNA extraction process in real-time PCR assays

#### Applications

- Easy validation of RNA extraction protocols
- Minimal interference with sample detection
- Includes a ready-to-use reaction mix for easy setup
- Suitable for use with blood, urine and sputum starting samples

The RT-qPCR Extraction Control enables users of diagnostic assays to validate both their extraction and RT-qPCR. Cells of a known concentration, containing the Internal Control RNA sequence are spiked into the sample tissue and RNA from the sample tissue and the RT-qPCR Extraction Control is simultaneously extracted.

Signal derived from the Internal Control RNA confirms the success of the extraction step and, as a known concentration of cells are added, RT-qPCR Extraction Control also monitors co-purification of PCR inhibitors that may cause biased or false amplification patterns.

#### Components

Reagent	500 Reactions
Internal Control RNA	5 x 200 μL
Control Mix	5 x 100 μL
50 mM MgCl <sub>2</sub>	1 x 1.2 mL

#### **Recommended Protocol**

Color coding	Internal	Control	Control	50 mM
	Control RNA	Mix 560	Mix 670	MgCl₂
Cap Colors	Purp <b>l</b> e	Yellow	Brown	Blue

All steps should be carried out at room temperature unless otherwise stated. Conditions may vary depending on the assay and may need optimization.

#### **Extraction step**

- 1. Briefly spin down all tubes before opening.
- 2. Standard Protocol:
  - i) Spike 2 µL of RT-qPCR Extraction Control (REC) into each sample
  - ii) Follow the manufacturer's protocol for total RNA extraction
  - iii) Elute total RNA in a volume of 100  $\mu L$
- 3. Use 5  $\mu$ L of the elution volume for a 20  $\mu$ L PCR reaction. For example: 2  $\mu$ L REC spiked into sample, Total sample RNA extracted and eluted in 100  $\mu$ L, 5  $\mu$ L RNA template is used for a 20  $\mu$ L reaction volume.

Note: This ratio (REC:Elution Vol:RNA template) must be maintained to avoid RNA Extraction Control failure

#### Post-extraction setup and analysis

The following real-time RT-PCR setup is recommended when the REC is to be used with the following: SensiFAST™ Probe One-Step No-ROX Kit

SensiFAST™ Probe One-Step Lo-ROX Kit SensiFAST™ Probe One-Step Hi-ROX Kit 1. Real-Time RT-PCR set up for SensiFAST Probe One-Step No-ROX Kit.

Component	Supplied	Volume
2x SensiFAST™ Probe One-Step No-ROX Mix*	No	10 µL
Target Primer/Probe mix	No	ΧμL
Extracted RNA template	No	ΧμL
Control Mix**	Yes	0.8 µL
50 mM MgCl₂	Yes	1.2 µL
Reverse transcriptase	No	0.2 μL
RiboSafe RNase inhibitor	No	0.4 µL
Total Volume (for 1 reaction)		20 µL

\* This also applies to any commercial real-time RT-PCR mix with a standard MgCl<sub>2</sub> concentration of 3 mM.

\*\* Vortex Control Mix tube before making up the mastermix.

Cycles	Temperature	Duration	Notes
1	42 °C	10-20 min	Reverse transcription
1	95 °C	3 min	Activation
	95 °C	10 s	Denaturation
30-40	60 °C†	30 s-45 s	Annealing/Extension/ Acquisition

2. Recommended reverse transcription and PCR cycling conditions.

<sup>†</sup> The standard annealing temperature is 60 °C, but may have to be optimized by the user, particularly if using an alternative commercial real-time RT-PCR mix



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Result	Target	REC	Interpretation
1	+	+	Target(s) and internal control RNA detected
2	-	+	Target(s) not detected, internal control RNA detected, indicates a successful extraction and real-time RT-PCR reaction
3	-	-	Invalid result: Target(s) and internal control RNA not detected, repeat test
4	+	-	Invalid result: Internal control not detected, repeat test

Note:

a) Validation of multiplex PCR should be performed prior to high throughput processes

b) The negative control reaction should contain all components required for amplification of sample RNA, including REC

c) A negative control ensures no cross-reactivity with the user-assay and  $\ensuremath{\mathsf{REC}}$ 

## Troubleshooting

Problem	Possible Cause	Recommendation	
Invalid Result or Internal Control failure	Not enough RNA template	The correct proportions are as follows: 2 $\mu L$ REC per clinical sample and an elution volume of 100 $\mu L$ . Check that the correct amount of extracted RNA template has been added to the reaction.	
	Real-time RT-PCR mix not compatible	The REC system requires extra magnesium, adjust final concentration to 6 mM final.	
	RNA contained an RT inhibitor	Remove inhibitors, such as SDS, EDTA, formamide and pyrophosphate, by ethanol precipitation of RNA, including a 70% ethanol wash step.	
Real-time RT-PCR failure*	Reaction conditions not optimal	Increase the primer annealing step from 30 s up to 45 s. Increase the reverse transcription step from 10 min up to 20 min.	
	RNA degraded	Analyze RNA on a denaturing gel to verify integrity. Ensure that all reagents are RNase-free.	
Poor specificity in	Primer dimers	Redesign primers to prevent self-annealing. Set up reactions on ice.	
real-time RT-PCR	Genomic DNA contamination	Treat RNA with DNase I and re-purify. If possible, use intron-spanning primers in real-time PCR.	
Significant shift in Ct	Inefficient extraction	Alter extraction protocol	

\* Shift in Ct or decrease in the fluorescence level (RFU) in the REC signal compared to the expected Ct or normalized fluorescence level

## **Technical Support:**

If the troubleshooting guide does not solve the difficulty you are experiencing, please contact Technical Support with details of reaction setup, cycling conditions and relevant data.

Email: tech@meridianlifescience.com

# **Associated Products**

Product	Pack size	Cat. No.
ISOLATE II RNA Mini Kit	10 Preps	BIO-52071
ISOLATE II RNA Plant Kit	10 Preps	BIO-52076
SensiFAST™ Probe One-Step Hi-ROX Kit	500 reactions	BIO-77005
SensiFAST™ Probe One-Step Lo-ROX Kit	500 reactions	BIO-78005
SensiFAST™ Probe One-Step No-ROX Kit	500 reactions	BIO-76005

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