# **Associated products**

Product	Description	Pack Size	Cat No.
ISOLATE II Genomic DNA Kit	Rapid isolation of high-quality genomic DNA from many different starting material	10 Preps 50 Preps 250 Preps	BIO-52065 BIO-52066 BIO-52067
ISOLATE II Plant DNA Kit	Rapid isolation of high-quality genomic DNA from a wide variety of plant species	10 Preps 50 Preps 250 Preps	BIO-52068 BIO-52069 BIO-52070
ISOLATE II RNA Mini Kit	Isolation of high-yield and extremely pure total RNA from a variety of samples	10 Preps 50 Preps 250 Preps	BIO-52071 BIO-52072 BIO-52073
ISOLATE II RNA Plant Kit	Plant Kit Isolation of high-yield and extremely pure total RNA from a wide variety of plant species		BIO-52076 BIO-52077
TRIsure™	Quick isolation of high-quality RNA from a variety of sources for subsequent use in cDNA synthesis	100 mL 200 mL	BIO-38032 BIO-38033
SensiFAST cDNA Synthesis Kit	Fully optimized to generate maximum yields of full-length and low abundance cDNA from RNA	50 Reactions 250 Reactions	BIO-65053 BIO-65054
Agarose Molecular biology grade agarose		100 g 500 g	BIO-41026 BIO-41025

# **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: mbi.tech@meridianlifescience.com

#### Licensing information

Purchase of this product conveys a licence from Life Technologies to use this SYBR® containing reagent in an end-user RUO assay. Parties wishing to incorporate this SYBR® containing reagent into a downstream kit, should contact Life Technologies for SYBR® Licencing information

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Storage and stability: The SensiFAST SYBR $^{\otimes}$  No-ROX One-Step Kit is shipped on dry/blue ice. All kit components should be stored at -20°C upon receipt. Excessive freeze/thawing is not recommended.

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

#### Quality control:

The SensiFAST SYBR® No-ROX One-Step Kit and its components are extensively tested for activity, processivity, efficiency, heat activation, sensitivity, absence of nuclease contamination and absence of nucleic acid contamination.

#### Safety precautions:

Please refer to the material safety data sheet for further information

For research or further manufacturing use only.

### Trademarks:

SensiFAST (Bioline Reagents Ltd), SYBR (Molecular Probes), iCycler MyiQ5, Opticon, Chromo4, Miniopticon, (Bio-Rad), LightCycler (Roche), StepOne (ABI), SmartCycler (CEPheid), RotorGene (Corbett), RealPlex (Eppendorf), Quantica (Techne), MX4000 (Stratagene).

# SensiFAST™ SYBR® No-ROX One-Step Kit

Shipping: On dry/blue ice Catalog numbers

Batch No.: See vial BIO-72001: 100 x 20 μL reactions: 1 x 1 mL

Concentration: see vial BIO-72005: 500 x 20 μL reactions: 5 x 1 mL



# Description

The SensiFAST™ SYBR® No-ROX One-Step Kit has been formulated for highly reproducible first-strand cDNA synthesis and subsequent real-time PCR (qPCR) in a single tube. A combination of the latest advances in buffer chemistry together with a reverse transcriptase and hot-start DNA polymerase system, ensures that SensiFAST SYBR® No-ROX One-Step Kit produces fast, highly-specific and ultra-sensitive one-step RT-qPCR.

The SensiFAST SYBR® No-ROX One-Step Kit consists of a 2x SensiFAST SYBR® One-Step mix, as well as separate reverse transcriptase and RiboSafe RNase Inhibitor.

### Kit components

Reagent	100 x 20 μL reactions	500 x 20 μL reactions
SensiFAST™ SYBR <sup>®</sup> No-ROX One-Step mix (2x)	1 x 1 mL	5 x 1 mL
RiboSafe RNase Inhibitor	1 x 40 μL	1 x 200 μL
Reverse transcriptase	1 x 20 μL	1 x 100 μL
DEPC-H <sub>2</sub> O	1 x 1.8 mL	2 x 1.8 mL

#### Instrument compatibility

SensiFAST SYBR® No-ROX One-Step Kit is premixed with SYBR Green I dve and is compatible with real-time PCR instruments that do not need a passive reference signal for normalization of the data. The SensiFAST SYBR® No-ROX One-Step Kit is optimized for use on the real-time instruments listed in the following compatibility table:

Manufacturer	Model
Bio-Rad	Opticon™, Opticon2™, MiniOpticon, Chromo4™, CFX96, CFX384
Cepheid	SmartCycler™
Qiagen (Corbett )	Rotor-Gene™ 3000 & 6000
Eppendorf	Realplex
Roche	LightCycler <sup>®</sup> 480
Techne	Quantica <sup>®</sup>

#### General considerations

When handling RNA, it is important to use RNase-free plasticware and reagents. We also recommend performing RNA work in an RNase-free area. To help prevent any carry-over DNA contamination, we recommend that separate areas are maintained for reaction set-up, PCR amplification and any post-PCR gel analysis. It is essential that any tubes containing amplified PCR product are not opened in the PCR set-up area.

**Primers:** The sequence and concentration of the primers, as well as amplicon length, can be critical for specific amplification, yield and overall efficiency of any RT-qPCR. We strongly recommend taking the following points into consideration when designing and running your RT-qPCR:

- use primer-design software, such as Primer3 or visual OMP<sup>TM</sup> (http://frodo.wi.mit.edu/primer3/ and DNA Software, Inc. http:// dnasoftware.com/, respectively). Primers should have a melting temperature (Tm) of approximately 60 °C
- optimal amplicon length should be 80-200 bp, and should not exceed 400 bp
- final primer concentration of 400 nM is suitable for most SYBR®-Green based reactions, however to determine the optimal concentration we recommend titrating in the range 0.1 -1 uM
- use an equimolar primer concentration
- where possible, use intron-spanning primers to avoid amplification from genomic DNA

Template: It is important that the RNA template is intact and devoid of DNA or contaminating inhibitors of both reverse transcription and PCR. For high purity RNA, we recommend using the ISOLATE II RNA Mini Kit (BIO-52073). RNA stocks and dilutions should be made in DEPC-treated water to avoid any RNase-mediated degradation.

The recommended amount of template for one-step RT-qPCR is dependent upon the type of RNA used.

- total RNA: purified total RNA can be used in the range from 1 pg to 1 μg per 20 μL reaction
- mRNA: purified mRNA can be used from 0.01 pg per 20 μL reaction

 $MgCl_2$ : The  $MgCl_2$  concentration in the 1x reaction mix is 3 mM. In the majority of RT-qPCR conditions this is optimal for both the reverse transcriptase and the hot-start DNA polymerase. If necessary, we suggest titrating the  $MgCl_2$  to a maximum of 5mM.

**RT-PCR controls:** It is important to detect the presence of contaminating DNA that may affect the reliability of the data. Always include a no-RT control, by omitting the reverse transcriptase from the reaction.

### **Procedure**

Reaction mix composition: Prepare an RT-qPCR master mix. The volumes given below are based on a standard 20  $\mu L$  final reaction mix and can be scaled accordingly.

Reagent	Volume	Final concentration
2x SensiFAST™ SYBR® No-ROX One-Step Mix	10 μL	1x
10 μM Forward Primer	0.8 μL	400 nM
10 μM Reverse Primer	0.8 μL	400 nM
Reverse transcriptase	0.2 μL	-
RiboSafe RNase Inhibitor	0.4 μL	-
H <sub>2</sub> O	up to 16 μL	
Template	4 μL	
	20 μL Final	volume

Sensitivity testing and C<sub>t</sub>values: When comparing SensiFAST with a mix from another supplier we strongly recommend amplifying from a 10-fold template dilution series. Loss of detection at low template concentration is the only direct measurement of sensitivity. An early C<sub>t</sub> value is not an indication of good sensitivity, but rather an indication of speed.

Suggested RT-qPCR conditions: The following RT-qPCR conditions are suitable for the SensiFAST SYBR® No-ROX One-Step Kit with the majority of amplicons and real-time PCR instruments. However, the cycling conditions can be varied to suit different machine-specific protocols. SensiFAST SYBR® No-ROX One-Step Kit is compatible with either three-step or two-step cycling:

## 3-step cycling

Cycles	Temp.	Time	Notes
1	45 °C	10 min	Reverse transcription
1	95 °C	2 min	Polymerase activation
40	95 °C 60 °C 72 °C	5 s 10 s 5 s	Denaturation Annealing Extension (acquire at end of step)

# 2-step cycling

Cycles	Temp.	Time	Notes
1	45 °C	10 min	Reverse transcription
1	95 °C	2 min	Polymerase activation
40	95 °C 60 °C	5 s 20 s	Denaturation Annealing/extension (acquire at end of step)

**Optional analysis:** After the reaction has reached completion, refer to the instrument instructions for the option of melt-profile analysis.

# Troubleshooting guide

Problem	Possible Cause	Recommendation
	Activation time too short	Ensure SensiFAST SYBR® No-ROX One-Step mix is activated for a minimum of 2 min at 95 °C before cycling
	Error in protocol setup	Verify that correct reagent concentrations, volumes, dilutions and storage conditions have been used
	Suboptimal primer design	Use primer design software or validated assay. Test assay on a control template
	Incorrect concentration of primers	Use primer concentrations between 100 nM and 1 μM
No amplification trace  AND  No product on	Template degraded	Re-isolate your template from the sample material or use freshly prepared template dilution. We recommend using the ISOLATE RNA kits for template preparation and DEPC-treated water for resuspension or dilution of the template
agarose gel		Verify the integrity of RNA using agarose gel electrophoresis
	Primers degraded	Use newly synthesized primers
	Template contaminated with RT-qPCR inhibitors	Further dilute template before RT-qPCR or purify template and resuspend it in DEPC-treated water
	Template concentration too low	Increase concentration used
	Cycling conditions not optimal	Increase extension/annealing time, increase cycle number, reduce annealing temperature

### Troubleshooting guide (Continued)

Problem	Possible Cause	Recommendation
No amplification trace AND PCR product present on agarose gel	Error in instrument setup	Check that the acquisition settings are correct during cycling
	Inefficient reverse transcription	Extend reverse transcription time up to 20 min and/or increase the temperature up to 48 °C
	Suboptimal primer design	Redesign primers using appropriate software, or use validated primers
Non-specific	Primer concentration too high	Test dilution series of primer concentrations until primer-dimer/non-specific amplification products disappear
amplification product	Primer concentration too low	Use primer concentration between 100 nM and 1 µM
AND Primer-dimers	Primer annealing temperature too low	Increase PCR annealing temperature up to 65 °C or until primer-dimer/non-specific amplification products disappear
	Template concentration too low	Increase template concentration
	Template concentration too high	Reduce template concentration until non-specific products disappear
	Extension time too long	Reduce extension time to determine whether non-specific products are reduced
Variability between	Error in reaction set-up	Prepare large volume master mix
replicates	Air bubbles in reaction mix	Centrifuge reaction samples/plate prior to running on a real-time instrument
	Inefficient reverse transcription	Extend reverse transcription time up to 20 min and/or increase the temperature up to 48 °C
	Activation time too short	Ensure SensiFAST SYBR <sup>®</sup> No-ROX One-Step mix is activated for a minimum of 1 min at 95 °C before cycling
	Annealing temperature too high	Decrease annealing temperature in steps of 2 °C
Late	Extension time too short	Double extension time to determine whether the cycle threshold $(C_T)$ is affected
amplification trace	Template concentration too low	Increase concentration if possible
	Template is degraded	Re-isolate template from sample material or use freshly prepared template dilution
	Suboptimal primer design	Redesign primers using appropriate software, or use validated primers
	Primer concentration too low	Increase concentration of primers in 100 nM increments
	RNase contamination	Ensure RNase inhibitor is added before addition of template
	Extension time too short	Increase extension time
PCR efficiency below 90%	Primer concentration too low	Increase concentration of primers in 100 nM increments
	Suboptimal primer design	Redesign primers using appropriate software or use validated primers
PCR officioney	Template is degraded or contains PCR inhibitors	Re-isolate template from sample material, or use freshly prepared template dilution, or purify template and resuspend it in water
efficiency above 110%	Non-specific amplification and/or primer-dimers	Use 4% agarose gel electrophoresis to confirm presence of non-specific amplification products. See above for preventing/removing non-specific products