BIÒSCIENCES[™] qScript[™] One-Step SYBR[®] Green qRT-PCR Kit for iQ[™]

Cat. No. 95086-050 95086-200 Size: 50 x 50-µL reactions 200 x 50-µL reactions Store at -20°C

Description

The qScript One-Step SYBR Green qRT-PCR Kit for iQ is a convenient and highly sensitive solution for reverse transcription quantitative PCR (RT-qPCR) of RNA templates using SYBR Green I dye detection and gene-specific primers on Bio-Rad iCycler iQ[®] systems. cDNA synthesis and PCR amplification are carried out in the same tube without opening between procedures. The system has been optimized to deliver maximum RT-qPCR efficiency, sensitivity, and specificity. The proprietary reaction buffer has been specifically formulated to maximize activities of both reverse transcriptase and Taq DNA polymerase while minimizing the potential for primer-dimer and other non-specific PCR artifacts. The kit is compatible with both fast and standard qPCR cycling protocols. Highly specific amplification is essential for successful RT-qPCR with SYBR Green I technology, since this dye binds to any dsDNA generated during amplification. AccuStart[™] Taq DNA polymerase contains monoclonal antibodies that bind to the polymerase and keep it inactive prior to the initial PCR denaturation step. Upon heat activation at 95°C, the antibodies denature irreversibly, releasing fully active, unmodified Taq DNA polymerase.

Instrument Compatibility

Different real-time PCR systems employ different strategies for the normalization of fluorescent signals and correction of well-to-well optical variations. It is critical to match the appropriate qPCR reagent and internal reference dye to your specific instrument. The qScript Custom One-Step SYBR Green qRT-PCR Kit for iQ contains fluorescein for experimental plate or dynamic well factor collection on iCycler iQ, $iQ^{TM}5$, or $MyiQ^{TM}$ real-time detection systems. Inclusion of fluorescein does not affect the reaction efficiency or sensitivity of detection. Please visit our web site at <u>www.quantabio.com</u> to find the optimal kit for your instrument platform.

Com	ponents
-----	---------

Reagent	Description	95086-050	95086-200
qScript One-Step Reverse Transcriptase	Optimized 50X formulation of recombinant MMLV reverse transcriptase for one- step RT-PCR.	1 x 50 µL	1 x 200 µL
One-Step SYBR Green Master Mix for iQ (2X)	2X reaction buffer containing dNTPs, magnesium chloride, AccuStart Taq DNA polymerase, stabilizers, fluorescein, and SYBR Green I dye	1 x 1.25 mL	4 x 1.25 mL
Nuclease-free water		1 x 1.5 mL	4 x 1.5 mL

Storage and Stability

Kit components are stable for one year when stored in a constant temperature freezer at -20° C, protected from light. For convenience, the One-Step SYBR Green Master Mix for iQ may be stored unfrozen at +2 to $+8^{\circ}$ C for up to 4 months.

Guidelines for One-Step SYBR Green qRT-PCR

- Primer design is critical for successful one-step RT-qPCR with SYBR Green. The use of software tools for PCR primer design and RNA secondary structure analysis can aide in the design of specific and efficient primers for one-step RT-qPCR. Primers should be designed according to standard qPCR guidelines with a length of 18 25 nucleotides and a GC content of 40-65%. Avoid internal secondary structure, and complementation at 3' ends within each primer and primer pair. 3'-end terminal stability should be kept low to maximize primer specificity (3'-pentamer ΔG° > -8.0 kcal/mol or have no more than 2 to 3 Cs or Gs in the last 5 bases).
- Regions of RNA secondary structure should be avoided as this can interfere with annealing of the reverse primer for cDNA synthesis and/or impede procession of the reverse transcriptase. Programs for RNA structure prediction, such as the mfold web server (<u>http://mfold.bioinfo.rpi.edu/</u>), are useful for selecting regions of relaxed RNA structure for qRT-PCR primer design.
- Ideally, primer Tm should be between 58 and 60°C for a typical 2-step qPCR cycling protocol. Estimation of primer Tm varies widely with different methods and analysis parameters. We recommend using a program that calculates Tm based on nearest-neighbor thermodynamic models at 50 mM monovalent salt and 50 nM primer concentration. Primers with melting temperatures outside of this range may require optimization of PCR cycling conditions.
- PCR product size should be between 70- 200 bp. Ideally, the amplified sequence should span intronic sequence to minimize the potential to amplify genomic DNA sequence. Design primers to anneal to exons that bracket intronic sequence or within exon / exon boundaries of the specific mRNA. NCBI's Primer-BLAST program (<u>http://www.ncbi.nlm.nih.gov/tools/primer-blast/index.cgi?LINK_LOC=BlastHomeAd</u>) can facilitate the design of RNA-specific primer sets. Control reactions that lack reverse transcriptase (minus RT) should always be included to verify that amplification signal is due to the presence of RNA target and not genomic DNA.
- A final concentration of 200 nM each primer is recommended as a general starting point. Optimal results may require titration of primer concentration between 100 and 500 nM. PCR efficiency is often improved with higher primer concentration (300 to 500 nM). In some cases, higher concentration of the reverse primer alone may improve RT-PCR efficiency without compromising specificity. We highly recommend including a post PCR dissociation analysis step (melt curve) to distinguish specific from non-specific amplification product(s) (i.e. primer-dimer).
- Thaw all components, except the qScript One-Step RT, at room temperature. Mix by gently vortexing, then centrifuge to collect contents to the bottom of the tube before using. Place all components on ice after thawing.
- To maximize assay specificity and sensitivity reactions should be assembled on ice and kept cold until placed in your real-time PCR system. Centrifugation steps should be carried out in a refrigerated centrifuge. AccuStart Taq DNA polymerase is inactive prior to high temperature activation; however, reverse transcriptases are active at lower temperatures and can use single strand DNA as a template.

Guidelines for One-Step SYBR Green qRT-PCR continued:

- First-strand synthesis can be carried out between 42°C and 52°C. Optimal results are generally obtained with a 5-minute incubation at 50°C. We recommend a 2-5 minute incubation at 95°C to fully inactivate the RT prior to PCR cycling.
- Preparation of a reaction cocktail is recommended to reduce pipetting errors and maximize assay precision. Assemble the reaction cocktail with all
 required components except RNA template and dispense equal aliquots into each reaction tube. Add RNA to each reaction as the final step.
 Addition of sample as 5 to 10-µL volumes will improve assay precision.
- Suggested input quantities of template are: 1 pg to 100 ng total RNA; 10 fg to 100 ng poly A(+) RNA; 10 to 1x10⁸ copies viral RNA.
- After sealing each reaction, vortex gently to mix contents. Centrifuge briefly to collect components at the bottom of the reaction tube.

Reaction Assembly		
Component	Volume for 50-µL rxn.	Final Concentration
One-Step SYBR Green Master Mix for iQ (2X)	25 μL	1X
Forward primer	Variable	200 – 300 nM
Reverse primer	Variable	200 – 300 nM
Nuclease-free water	Variable	
RNA template	5 — 10 μL	Variable
qScript One-Step RT *	<u>1 µL</u>	1X
Final Volume (µL)	50 µL	

Note: Reaction volume can be scaled from 5 to 50 µL depending on the reaction plate (i.e. 384-well vs. 96-well) and qPCR system. Scale all component volumes proportionally. * Omit addition of qScript One-Step RT in minus RT control reactions.

Reaction Protocol

Incubate the complete reaction mix in a real-time thermal detection system as follows:

	Fast qPCR Cycling	Standard qPCR Cycling	3-Step PCR Cycling
cDNA Synthesis	50°C, 5 min	48 – 50°C, 10 min	48 – 50°C, 10 min
Taq Activation	95°C, 2 min	95°C, 5 min	95°C, 5 min
PCR cycling (30 - 45 cycles)	95°C, 3s	95°C, 10s	95°C, 10s
	60°C, 30s (data collection)	60°C, 30s (data collection)	55 – 65°C, 20s
			68 – 72°C, 30 to 60s (data collection)
Melt Curve (dissociation stage):	See instrument instructions	See instrument instructions	See instrument instructions

Optimal cycling conditions will vary for different primer sets. A 3-step cycling protocol may improve assay specificity with some primer sets.

Quality Control

Kit components are free of contaminating DNase and RNase. The qScript One-Step SYBR Green qRT-PCR Kit for iQ is functionally tested in RT-qPCR. Kinetic analysis must demonstrate linear resolution over six orders of dynamic range ($r^2 > 0.995$) and an RT-PCR efficiency > 90%

Limited Label Licenses

This product is provided under an agreement between Molecular Probes, Inc. (a wholly owned subsidiary of Invitrogen Corporation) and Quanta Biosciences, Inc., and the manufacture, use, sale or import of this product is subject to one or more of U.S. Patent Nos. 5,436,134; 5,658,751 and corresponding international equivalents, owned by Molecular Probes. The purchase of this product conveys to the buyer the non-transferable right to use the purchased amount of the product and components of the product in research conducted by the buyer, where such research does not include testing, analysis or screening services for any third party in return for compensation on a per test basis. The buyer cannot sell or otherwise transfer (a) this product (b) its components or (c) materials made using this product or its components to a third party or otherwise use this product or its components or materials made using this product or its components for Commercial Purposes. Commercial Purposes means any activity by a party for consideration and may include, but is not limited to: (1) use of the product or its components in manufacturing; (2) use of the product or its components or not such product or its components are resold for use in research. For information on purchasing a license to this product for purposes; or (4) resale of the product or its components, whether or not such product or its components are resold for use in research. For information on purchasing a license to this product for purposes other than research, contact Molecular Probes, Inc., Business Development, 29851

Use of this product is covered by one or more of the following US patents and corresponding patent claims outside the US: 5,994,056 and 6,171,785. The purchase of this product includes a limited, nontransferable immunity from suit under the foregoing patent claims for using only this amount of product for the purchaser's own internal research. No right under any other patent claim and no right to perform commercial services of any kind, including without limitation reporting the results of purchaser's activities for a fee or other commercial consideration, is conveyed expressly, by implication, or by estoppel. This product is for research use only. Diagnostic uses under Roche patents require a separate license from Roche. Further information on purchasing licenses may be obtained by contacting the Director of Licensing, Applied Biosystems, 850 Lincoln Centre Drive, Foster City, California 94404, USA.

The purchase of this product includes a limited, non-transferable right to use the purchased amount of the product to perform Applied Biosystems' patented Passive Reference Method for the purchaser's own internal research. No right under any other patent claim and no right to perform commercial services of any kind, including without limitation reporting the results of purchaser's activities for a fee or other commercial consideration, is conveyed expressly, by implication, or by estoppel. This product is for research use only. For information about these rights or on obtaining additional rights, please contact outlicensing@lifetech.com or Out Licensing, Life Technologies, 5791 Van Allen Way, Carlsbad, California 92008.

The purchase of this product includes a limited, non-transferable license for all fields other than human or veterinary in vitro diagnostics under specific claims of U.S. Patent Nos. 6,174,670, 6,569,627 and 5,871,908, owned by the University of Utah Research Foundation or Evotec Biosystems GmbH and licensed to Idaho Technology, Inc. and Roche Diagnostics GmbH, to use only the enclosed amount of product according to the specified protocols. No right is conveyed, expressly, by implication, or by estoppel, to use any instrument or system under any claim of U.S. Patent Nos. 6,174,670, 6,569,627 and 5,871,908, other than for the amount of product contained herein.

Licensed to Quanta BioSciences, under U.S. Patent Nos. 5,338,671, 5,587,287, and foreign equivalents for use in research only.

qScript and AccuStart are trademarks of Quanta BioSciences Inc. SYBR is a registered trademark of Molecular Probes, Inc. iCycler iQ, iQ5, and MyiQ, are trademarks of Bio-Rad Laboratories.

©2009 Quanta BioSciences, Inc. All rights reserved. For research use only. Not intended for any animal or human therapeutic or diagnostic use.