

For life science research only.  
Not for use in diagnostic procedures.



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# LightCycler<sup>®</sup> DNA Master SYBR Green I

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 **Version 11**

Content version: April 2011

Easy-to-use Reaction Mix for PCR using the LightCycler<sup>®</sup> Carousel-Based System


**Cat. No. 12 015 099 001**

Kit for 96 reactions

**Cat. No. 12 158 817 001**

Kit for 480 reactions

**Store the kit at – 15 to –25°C**

 Keep LightCycler<sup>®</sup> DNA Master SYBR Green I (vial 1, green cap) away from light!

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# 1. What this Product Does

**Number of Tests** The kit is designed for 96 or 480 reactions (depending on pack size), with a final reaction volume of 20  $\mu$ l each.

## Kit Contents

Vial/Cap	Label	Contents/Function
		a) Cat. No. 12 015 099 001 (96 reactions) b) Cat. No. 12 158 817 001 (480 reactions)
1 green cap	LightCycler® DNA Master SYBR Green I, 10× conc.	a) 3 vials, 64 $\mu$ l each b) 15 vials, 64 $\mu$ l each ▪ Ready-to-use reaction mix for PCR ▪ Contains Taq DNA Polymerase, reaction buffer, dNTP mix (with dUTP instead of dTTP), SYBR Green I dye and 10 mM MgCl <sub>2</sub> .
2 blue cap	MgCl <sub>2</sub> stock solution, 25 mM	a) 1 vial, 1 ml b) 2 vials, 1 ml each ▪ To adjust MgCl <sub>2</sub> concentration.
3 colorless cap	H <sub>2</sub> O, PCR grade	a) 2 vials, 1 ml each b) 7 vials, 1 ml each ▪ To adjust the final reaction volume.

## Storage and Stability

Store the kit at –15 to –25°C until the expiration date printed on the label.

⚠ Keep the LightCycler® DNA Master SYBR Green I (vial 1, green cap) away from light!

🧊 The kit is shipped on dry ice.

Once the kit is opened, store the kit components as described in the following table:

Vial	Label	Storage
1 green cap	LightCycler® DNA Master SYBR Green I, 10× conc.	▪ Store at –15 to –25°C ▪ After thawing, store at +2 to +8°C for a maximum of two weeks. ▪ <b>Avoid repeated freezing and thawing!</b> ▪ <b>Keep vial 1 away from light!</b>
2 blue cap	MgCl <sub>2</sub> stock solution, 25 mM	Store at –15 to –25°C
3 colorless cap	H <sub>2</sub> O, PCR grade	

**Additional Equipment and Reagents Required**

Additional reagents and equipment required to perform PCR reactions with the LightCycler® DNA Master SYBR Green I, using the LightCycler® Carousel-Based System include:

- LightCycler® Carousel-Based System\* (LightCycler® 2.0 Instrument\*, or LightCycler® 1.5 Instrument\*, or an instrument version below)
- LightCycler® Capillaries\*
- Standard benchtop microcentrifuge, containing a rotor for 2.0 ml reaction tubes

Ⓢ The LightCycler® Carousel-Based System provides Centrifuge Adapters that enable LightCycler® Capillaries to be centrifuged in a standard microcentrifuge rotor.

or

- LC Carousel Centrifuge 2.0\* for use with the LightCycler® 2.0 Sample Carousel (20 µl; optional)

⚠ If you use a LightCycler® Instrument version below 2.0, you need in addition, the LC Carousel Centrifuge 2.0 Bucket 2.1\*. To adapt the LightCycler® 2.0 Sample Carousel (20 µl) to the former LC Carousel Centrifuge, you need the LC Carousel Centrifuge 2.0 Rotor Set\*.

- Nuclease-free, aerosol-resistant pipette tips
- Pipettes with disposable, positive-displacement tips
- Sterile reaction (Eppendorf) tubes for preparing master mixes and dilutions
- Uracil-DNA Glycosylase, heat-labile\* (optional<sup>†</sup>)

Ⓢ <sup>†</sup> For prevention of carry-over contamination; see section Related Procedures for details.

*\* available from Roche Applied Science; see Ordering Information for details.*

**Application**

LightCycler® DNA Master SYBR Green I is designed for research studies. When combined with the LightCycler® Carousel-Based System and suitable PCR primers, this kit enables very sensitive detection and quantification of defined DNA sequences. Furthermore, the kit can be used to perform two-step RT-PCR, in combination with a reverse transcription kit for cDNA synthesis\*.

In principle, the LightCycler® DNA Master SYBR Green I can be used for the amplification and detection of any DNA or cDNA target. However, you would need to optimize the detection protocol to the reaction conditions of the LightCycler® Carousel-Based System and design specific PCR primers for each target. Refer to the LightCycler® Operator’s Manual for general recommendations.

LightCycler® DNA Master SYBR Green I can also be used with Uracil-DNA Glycosylase, heat-labile, to prevent carry-over contamination during PCR.

⚠ The amplicon size should not exceed 1 kb in length. For optimal results, select a product length of 700 bp or less.

⚠ The performance of the kit described in this Instruction Manual is warranted only when it is used with the LightCycler® Carousel-Based System.

**Assay Time**

Procedure	Time
Optional: dilution of template DNA	5 min
PCR Setup	15 min
LightCycler® Carousel-Based System PCR run (incl. Melting Curve)	25 min
<b>Total assay time</b>	<b>45 min</b>

## 2. How to Use this Product

### 2.1 Before You Begin

- Sample Material**
- Use any template DNA (*e.g.*, genomic or plasmid DNA, cDNA) suitable for PCR in terms of purity, concentration and absence of inhibitors. For reproducible isolation of nucleic acids, use one of the following:
    - the MagNA Pure LC Instrument with a dedicated MagNA Pure LC reagent kit (for medium throughput automated isolation)
    - the MagNA Pure Compact Instrument with a dedicated MagNA Pure Compact reagent kit (for low throughput automated isolation)
    - a High Pure nucleic acid isolation kit (for manual isolation).

For further information, consult the Roche Applied Science Biochemicals catalog or home page: [www.roche-applied-science.com](http://www.roche-applied-science.com). See Ordering Information for selected products, recommended for the isolation of template DNA.

- Use up to 50 ng complex genomic DNA or  $10^1$  to  $10^{10}$  copies plasmid DNA.
- ⚠ Using a too high amount of template DNA may reduce the maximum fluorescence signal, by outcompeting the SYBR Green I dye.
- Ⓞ If you are using a non-purified cDNA sample from reverse transcription, especially if it contains high background concentrations of RNA and oligonucleotides, you can improve your results by using 2  $\mu$ l (or less) of that sample in the reaction.

**Primers** Use PCR primers at a final concentration of 0.2 to 1  $\mu$ M. The recommended starting concentration is 0.5  $\mu$ M each.

**MgCl<sub>2</sub>** To ensure specific and efficient amplification with the LightCycler® Carousel-Based System, you must optimize the MgCl<sub>2</sub> concentration for each target. The LightCycler® DNA Master SYBR Green I contains a MgCl<sub>2</sub> concentration of 1 mM (final concentration). The optimal concentration for PCR with the LightCycler® Carousel-Based System may vary from 1 to 5 mM. The table below gives the volumes of the MgCl<sub>2</sub> stock solution (vial 2, blue cap) that you must add to a 20  $\mu$ l reaction (final PCR volume), to increase the MgCl<sub>2</sub> concentration to the indicated values.

To reach a final Mg <sup>2+</sup> concentration (mM) of:	1	2	3	4	5
<b>Add this amount of 25 mM MgCl<sub>2</sub> stock solution (<math>\mu</math>l)</b>	0	0.8	1.6	2.4	3.2

The volume of water in the PCR reaction must be reduced, accordingly.

**Negative Control** Always run a negative control with the samples. To prepare a negative control, replace the template DNA with PCR-grade water (vial 3, colorless cap).

## 2.2 Experimental Protocol

### LightCycler® Carousel-Based System Protocol

The following procedure is optimized for use with the LightCycler® Carousel-Based System.

⚠ Program the LightCycler® Instrument before preparing the reaction mixes.

A LightCycler® Carousel-Based System protocol that uses the LightCycler® DNA Master SYBR Green I, contains the following programs:

- **Denaturation** of the template DNA
- **Amplification** of the target DNA
- **Melting Curve** for PCR product identification/amplicon analysis
- **Cooling** the rotor and the thermal chamber

For details on how to program the experimental protocol, see the LightCycler® Operator's Manual.

⚠ Set all other protocol parameters not listed in the table below to '0'.

The following table shows the PCR parameters that must be programmed for a LightCycler® Carousel-Based System PCR run with the LightCycler® DNA Master SYBR Green I.

Analysis Mode	Cycles	Segment	Target Temperature <sup>1)</sup>	Hold Time	Acquisition Mode
<b>Denaturation</b>					
None	1		95°C	30 s	none
<b>Amplification</b>					
Quantification	45	Denaturation	95°C	0 s	none
		Annealing	primer dependent <sup>2)</sup>	0 - 10 s <sup>4)</sup>	none
		Extension	72°C <sup>3)</sup>	= (amplicon [bp]/25) s <sup>5)</sup>	single
<b>Melting Curve</b>					
Melting Curves	1	Denaturation	95°C	0 s	none
		Annealing	65°C	15 s	none
		Melting	95°C Ramp Rate = 0.1°C/sec <sup>1)</sup>	0 s	continuous
<b>Cooling</b>					
None	1		40°C	30 s	none

<sup>1)</sup> Temperature Transition Rate/Slope/Ramp Rate is 20°C/sec, except where indicated.

<sup>2)</sup> For initial experiments, set the target temperature (*i.e.*, the primer annealing temperature) 5°C below the calculated primer  $T_m$ . Calculate the primer  $T_m$  according to the following formula, based on the nucleotide content of the primer:  $T_m = 2^\circ\text{C} (A+T) + 4^\circ\text{C} (G+C)$ .

<sup>3)</sup> If the primer annealing temperature is low (< +55°C), reduce the ramp rate to 2 to 5°C/s.

- 4) For typical primers, choose an incubation time of 0 to 10 s for the annealing step. To increase the specificity of primer binding, use an incubation time of < 5 s.
- 5) For greater precision in target quantification experiments, it can be advantageous (in some cases) to choose longer extension times for the amplification cycles.

## Fluorescence and Run Setup Parameters

Parameter	Setting	
<b>All LightCycler® Software Versions</b>		
Seek Temperature	30°C	
<b>LightCycler® Software prior to Version 3.5</b>		
Display Mode	Fluorescence channel F1	
Fluorescence Gains	Fluorimeter	Gain Value
	Channel 1 (F1)	3
	Channel 2 (F2)	1
	Channel 3 (F3)	1
<b>LightCycler® Software Version 3.5</b>		
Display Mode	Fluorescence channel F1	
Fluorescence Gains	not required	
	<p>Ⓞ In data created with LightCycler® Software Version 3.5, all fluorescence values are normalized to a fluorescence gain of “1”. This produces a different scale on the Y-axis than that obtained with previous LightCycler® Software versions. This difference does not affect the crossing points nor any calculated concentrations obtained.</p>	
<b>LightCycler® Software Version 4.1</b>		
Default Channel	Fluorescence channel 530	
Fluorescence Gains	not required	
“Max. Seek Pos.”	Enter the number of sample positions for which the Instrument should look.	
“Instrument Type”	<ul style="list-style-type: none"> <li>▪ “6 Ch.”: for LightCycler® 2.0 Instrument (selected by default)</li> <li>▪ “3 Ch.”: for LightCycler® 1.5 Instrument and instrument versions below</li> </ul>	
“Capillary Size”	Select “20 µl” as the capillary size for the experiment. ⚠ For the “6 Ch.” instrument type only.	

**Preparation of the PCR Mix**

Proceed as described below for a 20  $\mu$ l standard reaction.

⚠ Do not touch the surface of the capillaries. Always wear gloves when handling the capillaries.

- 1 Depending on the total number of reactions, place the required number of LightCycler® Capillaries in pre-cooled centrifuge adapters or in a LightCycler® Sample Carousel in a pre-cooled LC Carousel Centrifuge Bucket.
- 2
  - Thaw the solutions and for maximal recovery of contents, briefly spin vials in a microcentrifuge before opening.
  - Mix carefully by pipetting up and down and store on ice.
- 3 Prepare a 10 $\times$  conc. solution of the PCR primers.
  - Ⓞ If you are using the recommended final concentration of 0.5  $\mu$ M for each primer, the 10 $\times$  conc. solution would contain a 5  $\mu$ M concentration of each primer.

- 4 In a 1.5 ml reaction tube on ice, prepare the PCR Mix for one 20  $\mu$ l reaction, by adding the following components in the order mentioned below:

Component	Volume	Final conc.
H <sub>2</sub> O, PCR grade (vial 3, colorless cap)	x $\mu$ l	
MgCl <sub>2</sub> stock solution, 25 mM (vial 2, blue cap)	y $\mu$ l	Use concentration that is optimal for the target.
PCR Primer mix, 10 $\times$ conc.	2 $\mu$ l	0.2 to 1.0 $\mu$ M each (recommended conc. is 0.5 $\mu$ M)
LightCycler® DNA Master SYBR Green I, 10 $\times$ conc. (vial 1, green cap)	2 $\mu$ l	1 $\times$
<b>Total volume</b>	<b>18 <math>\mu</math>l</b>	

- Ⓞ To prepare the PCR Mix for more than one reaction, multiply the amount in the "Volume" column above by z, where z = the number of reactions to be run + one additional reaction.
- 5
  - Mix gently by pipetting up and down. Do not vortex.
  - Pipette 18  $\mu$ l PCR mix into each pre-cooled LightCycler® Capillary.
  - Add 2  $\mu$ l of the DNA template
  - Seal each capillary with a stopper.
- 6
  - Place the centrifuge adapters (containing the capillaries) into a standard benchtop microcentrifuge.
  - ⚠ Place the centrifuge adapters in a balanced arrangement within the centrifuge.
  - Centrifuge at 700  $\times$  g for 5 s (3,000 rpm in a standard benchtop microcentrifuge).
  - Alternatively, use the LC Carousel Centrifuge for spinning the capillaries.
- 7 Transfer the capillaries into the LightCycler® Sample Carousel and then into the LightCycler® Instrument.
- 8 Cycle the samples as described above.

## 2.3 Related Procedures

### Hot Start

If the reaction components are thoroughly mixed prior to the initial heat denaturation step, non-specific annealing and primer elongation events may occur. Conventional manual hot start or wax techniques cannot be used with the LightCycler® Carousel-Based System due to the small capillaries. However, you can run PCR applications that use the SYBR Green I detection format in hot start mode by using the LightCycler® FastStart DNA Master SYBR Green I or LightCycler® FastStart DNA Master<sup>PLUS</sup> SYBR Green I, which contain a chemically modified Taq DNA Polymerase, FastStart Taq DNA polymerase, that is activated by heat.

For details see the Roche Applied Science Biochemicals catalog or home page, [www.roche-applied-science.com](http://www.roche-applied-science.com).

### Prevention of Carry-Over Contamination

Uracil-DNA Glycosylase, heat-labile (UNG, heat-labile) is suitable for preventing carry-over contamination in PCR. This carry-over prevention technique involves incorporating deoxyuridine triphosphate (dUTP, a component of the reaction mixes of all LightCycler® reagent kits) into amplification products, then pretreating later PCR mixtures with UNG. If a dUTP-containing contaminant is present in the later PCRs, it will be cleaved by a combination of the UNG and the high temperatures of the initial denaturation step; it will not serve as a PCR template.


⚠ Use Uracil-DNA Glycosylase, heat-labile\* in combination with LightCycler® DNA Master SYBR Green I. Proceed as described in the package insert and/or in the table below, to prevent carry-over contamination. LightCycler® Uracil-DNA Glycosylase is to be used in combination with LightCycler® FastStart DNA Masters only.

- 
- ① Add 1  $\mu$ l UNG, heat-labile to the master mix per 20  $\mu$ l final reaction volume.
  - ② Add template DNA and incubate the completed reaction mixture for 5 min at room temperature (+15 to +25°C).
  - ③ Destroy any contaminating template and inactivate the UNG enzyme, by performing the initial denaturation step for 2 min at 95°C.
- 
- Ⓞ Since your target DNA template contains thymidine rather than uridine, it is not affected by this procedure.
- Ⓞ When performing Melting Curve analysis, the use of UNG may lower the melting temperature ( $T_m$ ) by approx. 1°C.
-

**Two Step RT-PCR** LightCycler® DNA Master SYBR Green I can also be used to perform two-step RT-PCR. In two-step RT-PCR, the reverse transcription of RNA into cDNA is separated from the other reaction steps and is performed outside the LightCycler® Carousel-Based System. Subsequent amplification and online monitoring is performed according to the standard LightCycler® Carousel-Based System procedure, using the cDNA as the starting sample material. One of the following reagents is required for reverse transcription of RNA into cDNA:

- Transcriptor Reverse Transcriptase\*
- Transcriptor First Strand cDNA Synthesis Kit\*
- First Strand cDNA Synthesis Kit for RT-PCR (AMV)\*

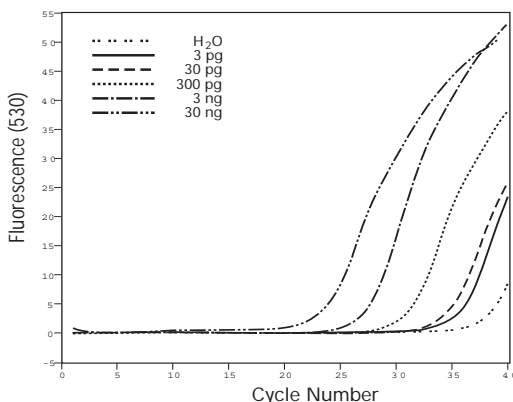
Synthesis of cDNA is performed according to the detailed instructions provided with the cDNA synthesis reagent.

 Do not use more than 8 µl of undiluted cDNA template per 20 µl final reaction volume, because greater amounts may inhibit PCR. For initial experiments, we recommend running undiluted, 1:10 diluted and 1:100 diluted cDNA template, in parallel to determine the optimal template amount.

### 3. Results

#### Quantification Analysis

The following amplification curves were obtained using the LightCycler® DNA Master SYBR Green I, in combination with the LightCycler® Control Kit DNA, targeting human  $\beta$ -globin gene. The fluorescence values versus cycle number are displayed. Thirty picograms (approx. 10 genome equivalents) of human genomic DNA can be reproducibly detected by amplifying in the LightCycler® Carousel-Based System and using the SYBR Green I detection format. Three picograms (approx. 1 haploid genome equivalent) are sporadically detected, due to statistical fluctuations.

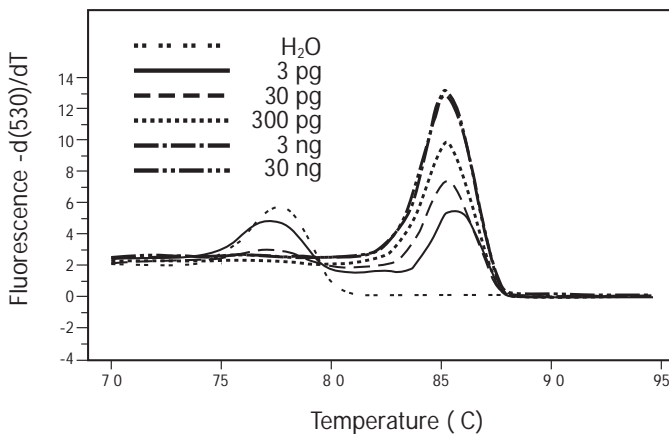


**Fig. 1:** Serially diluted samples containing 30 ng, 3 ng, 300 pg, 30 pg, or 3 pg human genomic DNA as starting template were amplified using the LightCycler® DNA Master SYBR Green I. As a negative control, template DNA was replaced by PCR-grade water.

## Melting Curve Analysis

Specificity of the amplified PCR product was assessed by performing a Melting Curve analysis. The resulting melting curves enable discrimination between primer-dimers and specific PCR product. The specific  $\beta$ -globin product melts at a higher temperature than the primer-dimers. The melting curves display the specific amplification of the  $\beta$ -globin gene when starting from 30 ng, 3 ng, 300 pg, 30 pg, or 3 pg human genomic DNA. Additionally, small amounts of PCR by-products are detected when starting from 30 pg and 3 pg human genomic DNA. In the negative control sample, only primer-dimers were amplified.

⚠️ Smaller reaction volumes may result in melting temperature variations.



**Fig. 2:** Melting Curve analysis of amplified samples containing 30 ng, 3 ng, 300 pg, 30 pg or 3 pg human genomic DNA as starting template. As a negative control, template DNA was replaced by PCR-grade water.

## 4. Troubleshooting

	Possible cause	Recommendation
Amplification reaches plateau phase before the program is complete.	Very high starting amount of nucleic acid	The program can be finished by clicking on the <b>End Program</b> button. The next cycle program will start automatically.
	The number of cycles is too high.	Reduce the number of cycles in the amplification program.
Log-linear phase of amplification just starts when the amplification program finishes.	Very low starting amount of nucleic acid	<ul style="list-style-type: none"> <li>Improve PCR conditions (<i>e.g.</i>, MgCl<sub>2</sub> concentration, primer concentration or design).</li> <li>Use higher amount of starting template.</li> <li>Repeat the run.</li> </ul>
	The number of cycles is too low.	<ul style="list-style-type: none"> <li>Increase the number of cycles in the amplification program.</li> <li>Use the <b>+10 cycles</b> button, to increase the number of cycles in the amplification program.</li> </ul>
No amplification occurs.	Wrong channel has been chosen to display amplification online.	Change the channel setting in the programming screen. (The data obtained up to this point will be saved.)
	Pipetting errors or omitted reagents	<ul style="list-style-type: none"> <li>Check for missing reagents.</li> <li>Titrate MgCl<sub>2</sub> concentration.</li> <li>Check for defective SYBR Green I dye.</li> </ul>
	Chosen gain settings are too low.	⚠ Optimize gain settings using the <b>Real Time Fluorimeter function</b> . Then repeat the run, using the optimal gain settings in the cycle programs.
	Scale of axes on graph are unsuitable for analysis.	Change the values for the x- and y-axis by double-clicking on the maximum and/or minimum values, then change to more suitable values.
	Measurements do not occur.	Check the amplification program. For SYBR Green I detection format, choose “single” as the acquisition mode at the end of the elongation phase.
	Amplicon length is >1 kb.	Do not use amplicons >1 kb. Optimal results are obtained with amplicons of 700 bp or less.
	Impure sample material inhibits the reaction.	<ul style="list-style-type: none"> <li>Do not use more than 8 to 10 µl of DNA per 20 µl PCR reaction mixture.</li> <li>Dilute the sample 1:10 and repeat the analysis.</li> <li>Repurify the nucleic acids to ensure removal of inhibitory agents.</li> </ul>

	Possible cause	Recommendation
Fluorescence intensity is too high and reaches overflow.	Unsuitable gain settings	<p>Gain settings cannot be changed during or after a run.</p> <p>Before repeating the run, use the <b>Real Time Fluorimeter</b> option to find suitable gain settings. For SYBR Green I, the background fluorescence at measuring temperature should not exceed 10.</p> <p>⚠ Use an extra sample for this procedure, so the dyes in your experimental samples will not be bleached. LightCycler® Software versions 3.5 and higher do not require a gain setting.</p>
Fluorescence intensity is too low.	Low concentration or deterioration of SYBR Green I dye in the reaction mixtures, due to unsuitable storage conditions.	<ul style="list-style-type: none"> <li>Store the SYBR Green I dye containing reagents at <math>-15</math> to <math>-25^{\circ}\text{C}</math> and keep them away from light.</li> <li>Avoid repeated freezing and thawing.</li> <li>After thawing, store the LightCycler® DNA Master SYBR Green I at <math>+2</math> to <math>+8^{\circ}\text{C}</math> for a maximum of two weeks and keep it away from light.</li> </ul>
	Reaction conditions are not optimized, leading to poor PCR efficiency.	<ul style="list-style-type: none"> <li>Titrate <math>\text{MgCl}_2</math> concentration.</li> <li>Primer concentration should be between 0.2 and <math>1.0\ \mu\text{M}</math></li> <li>Check annealing temperature of primers.</li> <li>Check experimental protocol.</li> <li>Always run a positive control along with your samples.</li> </ul>
Fluorescence intensity varies.	PCR mix is still in the upper part of the capillary. Air bubble is trapped in the capillary tip.	Repeat capillary centrifugation step.
Amplification curve reaches plateau at a lower signal level than the other samples.	Skin oils on the surface of the capillary tip	Always wear gloves when handling the capillaries.
	Starting amount of genomic DNA is too high; DNA captures SYBR Green I dye, leading to a high background signal. Insufficient amounts of SYBR Green I dye are left to monitor the increase of fluorescence signal during amplification.	<ul style="list-style-type: none"> <li>Do not use more than 50 ng of complex genomic DNA in a <math>20\ \mu\text{l}</math> reaction.</li> <li>Use the format of the HybProbe (which enables analysis of up to 500 ng DNA) instead of SYBR Green I.</li> </ul>
	SYBR Green I dye bleached.	Ensure the reagents containing the SYBR Green I dye are stored away from light. Avoid repeated freezing and thawing.

	Possible cause	Recommendation
Negative control samples are positive.	Contamination, or presence of primer-dimers.	<ul style="list-style-type: none"> <li>Remake all critical solutions.</li> <li>Pipette reagents on a clean bench.</li> <li>Close lid of the negative control reaction immediately after pipetting it.</li> <li>Use heat-labile UNG to eliminate carry-over contamination.</li> </ul>
Melting peak is very broad and peaks can not be differentiated.	°C to Average setting is too high.	Reduce the value of <b>°C to Average</b> (only applicable for LightCycler® Software versions prior to version 4.0).
Double melting peak appears for one product.	Two products of different length or GC content have been amplified (e.g., due to pseudogenes or mispriming).	<ul style="list-style-type: none"> <li>Check products on an agarose gel</li> <li>Elevate the reaction stringency by:                             <ul style="list-style-type: none"> <li>re-designing the primers,</li> <li>checking the annealing temperature,</li> <li>performing a “touch-down” PCR, or</li> <li>using HybProbe probes for better specificity.</li> </ul> </li> </ul>
Melting temperature of a product varies from experiment to experiment.	Variations in reaction mixture (e.g., salt concentration)	<ul style="list-style-type: none"> <li>Check purity of template solution.</li> <li>Reduce variations in parameters such as MgCl<sub>2</sub>, heat-labile UNG, hot-start antibody and program settings.</li> </ul>
Only a primer-dimer peak appears, with no specific PCR product peak; or very high primer-dimer peaks.	Primer-dimers have out-competed amplification of specific PCR product.	<ul style="list-style-type: none"> <li>Keep all samples at +2 to +8°C until the run is started.</li> <li>Keep the time between preparing the reaction mixture and starting the run as short as possible.</li> <li>Increase starting amount of DNA template.</li> <li>Titrate MgCl<sub>2</sub>.</li> <li>Increase annealing temperature, in order to enhance stringency.</li> </ul>
	Quality of the primers are poor.	<ul style="list-style-type: none"> <li>Purify primers more thoroughly.</li> <li>Use a hot start method.</li> </ul>
	Sequence of the primers are inappropriate.	Redesign primers.
Primer-dimer and product peaks are very close together.	Unusually high GC-content of PCR primers	<ul style="list-style-type: none"> <li>Redesign primers.</li> <li>Run melting curve at the lowest ramp rate (0.1 °C/sec with continuous measurement)</li> <li>Expand scale of the x-axis.</li> <li>Reduce the value of <b>°C to Average</b> (only applicable for LightCycler® Software versions prior to version 4.0).</li> </ul>
Very broad primer-dimer peak with multiple peaks	Heterogeneous primers with primer-dimer variations (e.g., concatamers, loops)	<ul style="list-style-type: none"> <li>Redesign primers.</li> <li>Use hot start method.</li> </ul>
One peak of the same height occurs in all samples.	Contamination in all samples	<ul style="list-style-type: none"> <li>Close capillaries during centrifugation step.</li> <li>Use fresh solutions.</li> </ul>

## 5. Additional Information on this Product

### 5.1 How this Product Works

LightCycler® DNA Master SYBR Green I is a ready-to-use PCR reaction mix. This product is designed specifically for real-time PCR assays in 20 µl glass capillaries, using the SYBR Green I detection format on the LightCycler® Carousel-Based System.

LightCycler® DNA Master SYBR Green I provides convenience, excellent performance and reproducibility, as well as minimizing contamination risk. All you must supply is template DNA, PCR primers and additional MgCl<sub>2</sub> (if necessary).

### Test Principle

Generation of PCR products can be detected by measurement of the SYBR Green I fluorescence signal. SYBR Green I intercalates into the DNA double helix. In solution, the unbound SYBR Green I dye exhibits very little fluorescence; however, fluorescence (wavelength, 530 nm) is greatly enhanced upon DNA-binding. Therefore, during PCR, the increase in SYBR Green I fluorescence is directly proportional to the amount of double-stranded DNA generated.

As SYBR Green I dye is very stable (only 6% of the activity is lost during 30 amplification cycles) and the LightCycler® Instruments' optical filter set matches the wavelengths of excitation and emission, it is the reagent of choice when measuring total DNA.

The basic steps of DNA detection by SYBR Green I, during real-time PCR on the LightCycler® Carousel-Based System are:

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- ① At the beginning of amplification, the reaction mixture contains the denatured DNA, the primers and the SYBR Green I dye. The unbound SYBR Green I dye molecules weakly fluoresce, producing a minimal background fluorescence signal, which is subtracted during computer analysis.

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  - ② After annealing of the primers, a few SYBR Green I dye molecules can bind to the double strand. DNA binding results in a dramatic increase of the SYBR Green I dye molecules to emit light upon excitation.

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  - ③ During elongation, more and more SYBR Green I dye molecules bind to the newly synthesized DNA. If the reaction is monitored continuously, an increase in fluorescence is viewed in real-time. Upon denaturation of the DNA for the next heating cycle, the SYBR Green I dye molecules are released and the fluorescence signal falls.

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  - ④ Fluorescence measurement at the end of the elongation step of every PCR cycle is performed, to monitor the increasing amount of amplified DNA.
-

To prove that only your desired PCR product has been amplified, you may perform a Melting Curve analysis after PCR. In Melting Curve analysis, the reaction mixture is slowly heated to +95°C, which causes melting of double-stranded DNA and a corresponding decrease in SYBR Green I fluorescence. The Instrument continuously monitors this fluorescence decrease and displays it as melting peaks. Each melting peak represents the characteristic melting temperature ( $T_m$ ) of a particular DNA product (where the DNA is 50% double-stranded and 50% single-stranded). The most important factors that determine the  $T_m$  of dsDNA are the length and the GC-content of that fragment. If PCR generated only one amplicon, Melting Curve analysis will show only one melting peak. If primer-dimers or other non-specific products are present, they will be shown as additional melting peaks. Checking the  $T_m$  of a PCR product can thus be compared with analyzing a PCR product by length in gel electrophoresis.

## 5.2 Quality Control

The LightCycler® DNA Master SYBR Green I is function tested using the LightCycler® Carousel-Based System.

### 5.3 Product Citations

- 1 Plachý, R. *et al.* (2005). McRAPD as a new approach to rapid and accurate identification of pathogenic yeasts. *J. Microbiol. Methods* **60**, 107-113.
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- 4 Hartmann, J. *et al.* (2004). Distinct Roles of G $\alpha_q$  and G $\alpha_{11}$  for Purkinje Cell Signaling and Motor Behavior. *J. Neurosci.* **24**, 5119-5130.
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- 7 Ohtani, S. *et al.* (2004). Quantitative analysis of *p53*-targeted gene expression and visualization of *p53* transcriptional activity following intratumoral administration of adenoviral *p53* *in vivo*. *Mol. Cancer Ther.* **3**, 93-100.
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- 9 Broberg, EK. *et al.* (2003). Low copy number detection of herpes simplex virus type 1 mRNA and mouse Th1 type cytokine mRNAs by Light Cycler quantitative real-time PCR. *J. Virol. Methods* **112**, 53-65.
- 10 Moeller, F. *et al.* (2003). New tools for quantifying and visualizing adoptively transferred cells in recipient mice. *J. Immunol. Methods* **282**, 73-82.
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- 13 Worm, J. *et al.* (2001). In-Tube DNA Methylation Profiling by Fluorescence Melting Curve Analysis. *Clin. Chem.* **47**, 1183-1189.

## 6. Supplementary Information

### 6.1 Conventions

**Text Conventions** To make information consistent and memorable, the following text conventions are used in this Instruction Manual:

Text Convention	Usage
Numbered stages labeled ①, ② etc.	Stages in a process that usually occur in the order listed.
Numbered instructions labeled ①, ② etc.	Steps in a procedure that must be performed in the order listed.
Asterisk *	Denotes a product available from Roche Applied Science.

### Symbols

In this Instruction Manual, the following symbols are used to highlight important information:

Symbol	Description
ⓘ	Information Note: Additional information about the current topic or procedure.
⚠	Important Note: Information critical to the success of the procedure or use of the product.

### Abbreviations

In this Instruction Manual, the following abbreviations are used:

Abbreviation	Meaning
dsDNA	double-stranded DNA
$T_m$	melting temperature
UNG	Uracil-DNA Glycosylase

### 6.2 Changes to Previous Version

Update of License Disclaimer

### 6.3 Ordering Information

Roche Applied Science offers a large selection of reagents and systems for life science research. For a complete overview of related products and manuals, please visit and bookmark our home page: [www.roche-applied-science.com](http://www.roche-applied-science.com) and our Special Interest Sites, including:

- Real-time PCR Systems (LightCycler<sup>®</sup> Carousel-Based System, LightCycler<sup>®</sup> 480 System, LightCycler<sup>®</sup> 1536 System, RealTime ready qPCR assays and Universal ProbelLibrary):  
<http://www.lightcycler.com>
- Automated Sample Preparation (MagNA Lyser Instrument, MagNA Pure Compact System, MagNA Pure LC Systems and MagNA Pure 96 System):  
<http://www.magnapure.com>

	<b>Product</b>	<b>Pack Size</b>	<b>Cat. No.</b>
Instrument and Accessories	LightCycler <sup>®</sup> 2.0 Instrument	1 instrument plus accessories	03 531 414 001
	LightCycler <sup>®</sup> 1.5 Instrument	1 instrument plus accessories	04 484 495 001
	LightCycler <sup>®</sup> Capillaries (20 µl)	1 pack (5 boxes, each with 96 capillaries and stoppers)	04 929 292 001
	LightCycler <sup>®</sup> Software 4.1	1 software package	04 898 915 001
	LC Carousel Centrifuge 2.0	1 centrifuge plus rotor and bucket	03 709 507 001 (115 V) 03 709 582 001 (230 V)
	LC Carousel Centrifuge 2.0 Rotor Set	1 rotor + 2 buckets	03 724 697 001
	LC Carousel Centrifuge 2.0 Bucket 2.1	1 bucket	03 724 689 001
	MagNA Pure LC 2.0 Instrument	1 instrument plus accessories	05 197 686 001
	MagNA Pure Compact Instrument	1 instrument plus accessories	03 731 146 001
	DNA Isolation Kits	MagNA Pure LC DNA Isolation Kit I	1 kit (192 isolations)
MagNA Pure LC DNA Isolation Kit II (Tissue)		1 kit (192 isolations)	03 186 229 001
MagNA Pure LC DNA Isolation Kit III (Bacteria, Fungi)		1 kit (192 isolations)	03 264 785 001
MagNA Pure LC DNA Isolation Kit - Large Volume		1 kit (96 - 288 isolations)	03 310 515 001
Total Nucleic Acid Isolation Kits	MagNA Pure LC Total Nucleic Acid Isolation Kit	1 kit (192 isolations)	03 038 505 001

	<b>Product</b>	<b>Pack Size</b>	<b>Cat. No.</b>	
RNA Isolation Kits	MagNA Pure LC Total Nucleic Acid Isolation Kit - Large Volume	1 kit (192 isolations)	03 264 793 001	
	MagNA Pure LC Total Nucleic Acid Isolation Kit - High Performance	1 kit (96 – 288 isolations)	05 323 738 001	
	MagNA Pure Compact Nucleic Acid Isolation Kit I	1 kit (32 isolations)	03 730 964 001	
	MagNA Pure Compact Nucleic Acid Isolation Kit I - Large Volume	1 kit (32 isolations)	03 730 972 001	
	MagNA Pure LC RNA Isolation Kit - High Performance	1 kit (192 reactions)	03 542 394 001	
	MagNA Pure LC RNA Isolation Kit III (Tissue)	1 kit (192 reactions)	03 330 591 001	
	MagNA Pure LC mRNA HS Kit <sup>1</sup>	1 kit (192 reactions)	03 267 393 001	
	MagNA Pure Compact RNA Isolation Kit	1 kit (32 reactions)	04 802 993 001	
	LightCycler <sup>®</sup> Kits for PCR	LightCycler <sup>®</sup> DNA Master HybProbe	1 kit (96 reactions)	12 015 102 001
		LightCycler <sup>®</sup> FastStart DNA Master HybProbe	1 kit (480 reactions)	12 158 825 001
LightCycler <sup>®</sup> FastStart DNA Master HybProbe		1 kit (96 reactions)	03 003 248 001	
LightCycler <sup>®</sup> FastStart DNA Master <sup>PLUS</sup> HybProbe		1 kit (480 reactions)	12 239 272 001	
LightCycler <sup>®</sup> FastStart DNA Master <sup>PLUS</sup> HybProbe		1 kit (96 reactions)	03 515 575 001	
LightCycler <sup>®</sup> FastStart DNA Master <sup>PLUS</sup> HybProbe		1 kit (480 reactions)	03 515 567 001	
LightCycler <sup>®</sup> FastStart DNA Master <sup>PLUS</sup> HybProbe		1 kit (384 reactions, 100 µl)	03 752 178 001	
LightCycler <sup>®</sup> FastStart DNA Master SYBR Green I		1 kit (96 reactions)	03 003 230 001	
LightCycler <sup>®</sup> FastStart DNA Master SYBR Green I		1 kit (480 reactions)	12 239 264 001	
LightCycler <sup>®</sup> FastStart DNA Master <sup>PLUS</sup> SYBR Green I		1 kit (96 reactions)	03 515 869 001	
LightCycler <sup>®</sup> FastStart DNA Master <sup>PLUS</sup> SYBR Green I	1 kit (480 reactions)	03 515 885 001		
LightCycler <sup>®</sup> FastStart DNA Master <sup>PLUS</sup> SYBR Green I	1 kit (384 reactions, 100 µl)	03 752 186 001		
cDNA Synthesis Reagents	Transcriptor Reverse Transcriptase	250 U	03 531 317 001	
	Transcriptor Reverse Transcriptase	500 U	03 531 295 001	
	Transcriptor Reverse Transcriptase	2,000 U	03 531 287 001	
	Transcriptor First Strand cDNA Synthesis Kit	1 kit (50 reactions, incl. 10 control reactions)	04 379 012 001	
	Transcriptor First Strand cDNA Synthesis Kit	1 kit (100 reactions)		
	Transcriptor First Strand cDNA Synthesis Kit	1 kit (200 reactions)	04 896 866 001	
Transcriptor First Strand cDNA Synthesis Kit for RT-PCR (AMV)		04 897 030 001		
Transcriptor First Strand cDNA Synthesis Kit for RT-PCR (AMV)	1 kit (30 reactions)	11 483 188 001		
Associated Kits and Reagents	Uracil-DNA Glycosylase, heat-labile	100 U	11 775 367 001	
	Uracil-DNA Glycosylase, heat-labile	500 U	11 775 375 001	

<b>Product</b>	<b>Pack Size</b>	<b>Cat. No.</b>
LightCycler® Control Kit DNA	1 kit (50 reactions)	12 158 833 001
High Pure PCR Template Preparation Kit	1 kit (100 purifications)	11 796 828 001
High Pure Plasmid Isolation Kit	1 kit (50 purifications)	11 754 777 001
	1 kit (250 purifications)	11 754 785 001

<sup>1</sup> the MagNA Pure LC mRNA HS Kit is only available for use with the MagNA Pure LC 1.0 Instrument (Cat. No. 12 236 931 001).

#### 6.4 Notice to Purchaser

NOTICE: This product may be subject to certain use restrictions. Before using this product please refer to the Online Technical Support page (<http://technical-support.roche.com>) and search under the product number or the product name, whether this product is subject to a license disclaimer containing use restrictions.

#### 6.5 Trademarks

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Exiqon and ProbeLibrary are registered trademarks of Exiqon A/S, Vedbaek, Denmark.

#### 6.6 Regulatory Disclaimer

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## Contact and Support

If you have questions or experience problems with this or any Roche Applied Science (RAS) product, please contact our Technical Support staff. Our scientists commit themselves to providing rapid and effective help.

We also want you to contact us if you have suggestions for enhancing RAS product performance or using our products in new or specialized ways. Such customer information has repeatedly proven invaluable to RAS and the worldwide research community.

To ask questions, solve problems, suggest enhancements or report new applications, please visit our **Online Technical Support Site at:**

**[www.roche-applied-science.com/support](http://www.roche-applied-science.com/support)**

To call, write, fax, or email us, visit the Roche Applied Science home page, [www.roche-applied-science.com](http://www.roche-applied-science.com), and select your home country.

Country-specific contact information will be displayed.

On the Roche Applied Science home page select **Printed Materials** to find:

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- our quarterly Biochemica Newsletter
- Material Safety Data Sheets
- Pack Inserts and Product Instructions

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