

LightMix[®] Kit *Trichomonas vaginalis* Cat.-No. 40-0610-32

Kit with reagents for the detection of *Trichomonas vaginalis* using the Roche Diagnostics LightCycler[®] 1.x / 2.0 / 480 II / Cobas[®] Z480 (open channel) Instruments.

Lyophilized mix of primers and probes for a total of 96 reactions with a final volume of 20 µl each.
Store protected from light at room temperature (18-25°C), do NOT freeze!

Instructions for use with the LightCycler[®] 1.x / 2.0 Instruments see pages 4-5
Instructions for use with the LightCycler[®] 480 / Cobas[®] Z480 Instruments see pages 6-7

1. Introduction

The LightMix[®] Kit for the detection of DNA from *Trichomonas vaginalis* provides a fast, easy and accurate system to identify this target in a nucleic acid extract. A second amplification reaction acts as internal control (IC).

This LightMix[®] Kit is tested on the LightCycler[®] 1.x / 2.0 / 480 II Instruments with Roche Diagnostics 'LightCycler[®] FastStart DNA Master HybProbe'.

2. Description

A 145 bp long fragment of the 'repeated DNA target'^{1,2} from the genome of *Trichomonas vaginalis* (*T. vaginalis*) is amplified with specific primers and detected with hybridization probes labeled with LightCycler[®] Red 640 (detected in channel 640).

An additional PCR product of 125 bp length is formed from the internal control (IC) DNA target. This control amplification does not interfere with the *T. vaginalis* specific reaction; amplification will usually fail in the presence of higher concentrated *T. vaginalis* samples (1,000 - 10,000 copies or higher) but will display an amplification signal in negative and low-concentrated samples. The probes are labeled with the dye LC690. Signals are recorded in channel 705 (660). The IC is supplied separately to allow running the assay with or without IC.

The use of a color compensation file generated with the TIB MOLBIOL 'LightMix[®] Kit - Color Compensation HybProbe' is a prerequisite to run the duplex or triplex reaction.

The supplied standard row allows to determine the linear range of the reaction and to estimate the quantity of the target sequence in unknown samples.

For use in LightCycler[®] 1.x Instruments with software version 3.5.3 read channel F2 instead of channel 640, channel F3 instead of channel 705 and channel F1 instead of channel 530 for detection. We recommend upgrading LightCycler[®] 1.x Instruments to software version 4.1.

¹ *Trichomonas vaginalis*: repeated DNA target for highly sensitive and specific polymerase chain reaction diagnosis. Kengne,P., Veas,F., Vidal,N., Rey,J.L. and Cuny,G. Cell. Mol. Biol. (Noisy-le-grand) 40 (6), 819-831 (1994)

² *Trichomonas vaginalis*: repeated DNA target for highly sensitive and specific polymerase chain reaction diagnosis. Kengne P, Veas F, Vidal N, Rey JL, Cuny G. Cell Mol Biol (Noisy-le-grand) 1994 Sep;40(6):819-31

3. Set Contents

- 3 Vials with **green** cap containing premixed primers and probes for each 32 PCR reactions
- 3 Vials with **white** cap containing the internal control (IC)
- 1 Standard row with 6 lyophilized standards *T. vaginalis* from 10¹ to 10⁶ target equivalents per rxn
- 1 Sealing foil for the standard row

4. Additional Reagents and items required

ColorCompensation HybProbe order n°40-0318-00	Roche Diagnostics Cat.-No. 05 997 704 001
LightCycler® FastStart DNA Master HybProbe	Cat.-No. 03 003 248 001
High Pure PCR Template Preparation Kit	Cat.-No. 11 796 828 001
LightCycler® Capillaries (20 µl) (LightCycler® 1.x / 2.0 Instruments)	Cat.-No. 04 929 292 001
LightCycler® 480 Multiwell Plate 384, white (LightCycler® 480 Instrument)	Cat.-No. 04 729 749 001
or LightCycler® 480 Multiwell Plate 96, white (LightCycler® 480 Instrument)	Cat.-No. 04 729 692 001

5. Product Characteristics

PCR results are obtained within 50 minutes (50 cycles and melting curve) with the LightCycler® 1.x / 2.0 Instruments and within 80 minutes (50 cycles and melting curve) with the LightCycler® 480 Instruments.

Sensitivity

These reagents detect 10 copies of *T. vaginalis* DNA using the Roche 'LightCycler® FastStart DNA Master HybProbe' with the LightCycler® 1.x / 2.0 / 480 II / Z480 Instruments (in an exemplary system, using cloned targets as reference).

Measuring range

The linear measuring range of the assay is 10² to 10⁶ copies of *T. vaginalis* DNA using the Roche 'LightCycler® FastStart DNA Master HybProbe' with the LightCycler® 1.x / 2.0 / 480 II / Z480 Instruments.

Storage and Stability

- Lyophilized reagents are stable for at least 6 months after shipment when stored protected from light at room temperature (18-25°C). See expiry date on the product label.
- **Do not freeze** lyophilized reagents.
- Dissolved reagents are stable for at least 10 days when stored protected from light and refrigerated (4°C).

6. Experimental Protocol

The following procedure was developed for use with the LightCycler® 1.x / 2.0 / 480 II Instruments. Start programming before preparing the solutions. See the LightCycler® operator's manual for details.

Sample material: Use aqueous nucleic acid preparations (e.g. 'High Pure PCR Template Prep. Kit').

Negative control: Always run at least one no-template control (NTC) - replace the template DNA with water.

Positive control: Run a positive control - replace the template DNA with the provided control DNA.

6.1. Preparation of parameter-specific reagents and reagents for the IC (32 reactions):

One reagent vial with a **green** cap contains primers and probes to run 32 reactions for for *T. vaginalis*. One reagent vial with a **white** cap contains primers, probes and DNA to run 32 reactions for the IC.

Add 66 µl PCR-grade water to each reagent vial, mix the solution (vortex) and spin down.

► **Use 2 µl** reagent for a 20 µl PCR reaction.

| This solution is stable at least five days when stored refrigerated at 4°C. Avoid prolonged exposure to light.

6.2. Preparation of the standard row

The target DNA is provided in 6 different quantities to yield from 10^1 to 10^6 target molecules in 5 µl once resolved. Start with the lowest concentrated standard (first tube from the extended lip). Use the pipette tip to punch a hole through the sealing foil. **Add 40 µl** PCR-grade water to each vial of the row. Mix the target DNA by pipetting the solution up and down 10 times.



► **Use 5 µl** standard for a 20 µl PCR reaction.

This standard solution is not long-term stable and will lose sensitivity under prolonged storage. Use only fresh prepared solutions as quantification references. Older solutions can be used as a qualitative standard (positive control). After adding the target DNA to the LightCycler® reaction mix use the provided sealing foil to close the vials in order to avoid changes in the concentration due to evaporation.

Please note that opening these vials may cause contaminations of the work-space (aerosol).

6.3. Preparation of the LightCycler® reaction mix

In a cooled reaction tube, prepare the reaction mix by multiplying each volume for a single reaction by the number of reactions to be cycled plus one additional reaction.

For use with the Roche FastStart Master		
Single reaction	Component	32 reactions
7.4 µl	water, PCR-grade (colorless cap, provided with the Roche Master kit)	236.8 µl
1.6 µl	Mg ²⁺ solution 25 mM (blue cap, provided with the Roche FastStart kit)	51.2 µl
2.0 µl	reagent mix (parameter specific reagents containing primers and probes, see 6.1.)	64.0 µl
2.0 µl	IC mix (IC reagents containing primers, probes and DNA, see 6.1.)	64.0 µl
2.0 µl	Roche Master (red cap, for preparation see Roche manual)	64.0 µl
15.0 µl	Volume of reaction mix	480 µl

Table 1

To include the internal control **add 2 µl** of the IC reagent per reaction to the reaction mix.

To run the assay without the internal control, substitute the 2 µl of IC with 2 µl PCR-grade water.

Mix gently, spin down and **transfer 15 µl** each of the reaction mix to a LightCycler® capillary (LightCycler® 1.x / 2.0 Instrument) or to a multiwell plate (LightCycler® 480 II Instrument).

Add 5 µl of sample or standard to each capillary or well for a final reaction volume of 20 µl.

Start run.

7. LightCycler® 1.x / 2.0 Instruments

7.1. Programming

The protocol consists of four program steps

- 1: Denaturation: sample denaturation and enzyme activation
- 2: Cycling: PCR-amplification of the target DNA
- 3: Melting: melting curve analysis for identification of the PCR product derived from the target DNA
- 4: Cooling: cooling the instrument

Program Step:	Denaturation	Cycling			Melting			Cooling
Parameter								
Analysis Mode	None	Quantification mode			Melting Curves mode			None
Cycles	1	50			1			1
Target [°C]	95	95	62	72	95	40	85	40
Hold [hh:mm:ss]	00:10:00	00:00:05	00:00:05	00:00:15	00:00:20	00:00:20	00:00:00	00:00:30
Ramp Rate [°C/s]	20	20	20	20	20	20	0.2	20
Sec Target [°C]	-	-	55	-	-	-	-	-
Step Size [°C]	-	-	0.5	-	-	-	-	-
Step Delay (Cycles)	-	-	1	-	-	-	-	-
Acquisition Mode	None	None	Single	None	None	None	Cont	None

Table 2

(Melting not relevant for detection)

7.2. Data Analysis

Switch the color compensation mode on. If this mode is not enabled run the color compensation program. Follow the instructions in the manual of "LightMix® Kit – Color Compensation 530/640/690".

Perform data analysis, as described in the LightCycler® Instrument operator's manual.

We recommend using the Second Derivative Maximum method (Automated (F" max)). The cycle number of the Crossing Point (Cp) of each sample is calculated automatically. The Fit Points method is more-error prone due to the user's influence.

View *T. vaginalis* data in channel 640, Quantification mode. The negative control (NTC) must show no signal. For the identification of the PCR product view *T. vaginalis* data in channel 640 Melting Curves mode.

If the internal control (IC) is used view IC data in channel 705, Quantification mode. The negative control and the low-concentrated *T. vaginalis* DNA samples (10 to 1,000 copies) should show an amplification curve for the IC with a Cp at approximately cycles 29- 31.

The provided standard row of cloned and purified DNA with concentrations in the range from 10⁶ copies/rxn to 10¹ copies/rxn of *T. vaginalis* should have Cp values between cycles 17 and 35 (Cp values calculated with Second Derivative Maximum method).

7.3. Sample Data – Typical Results

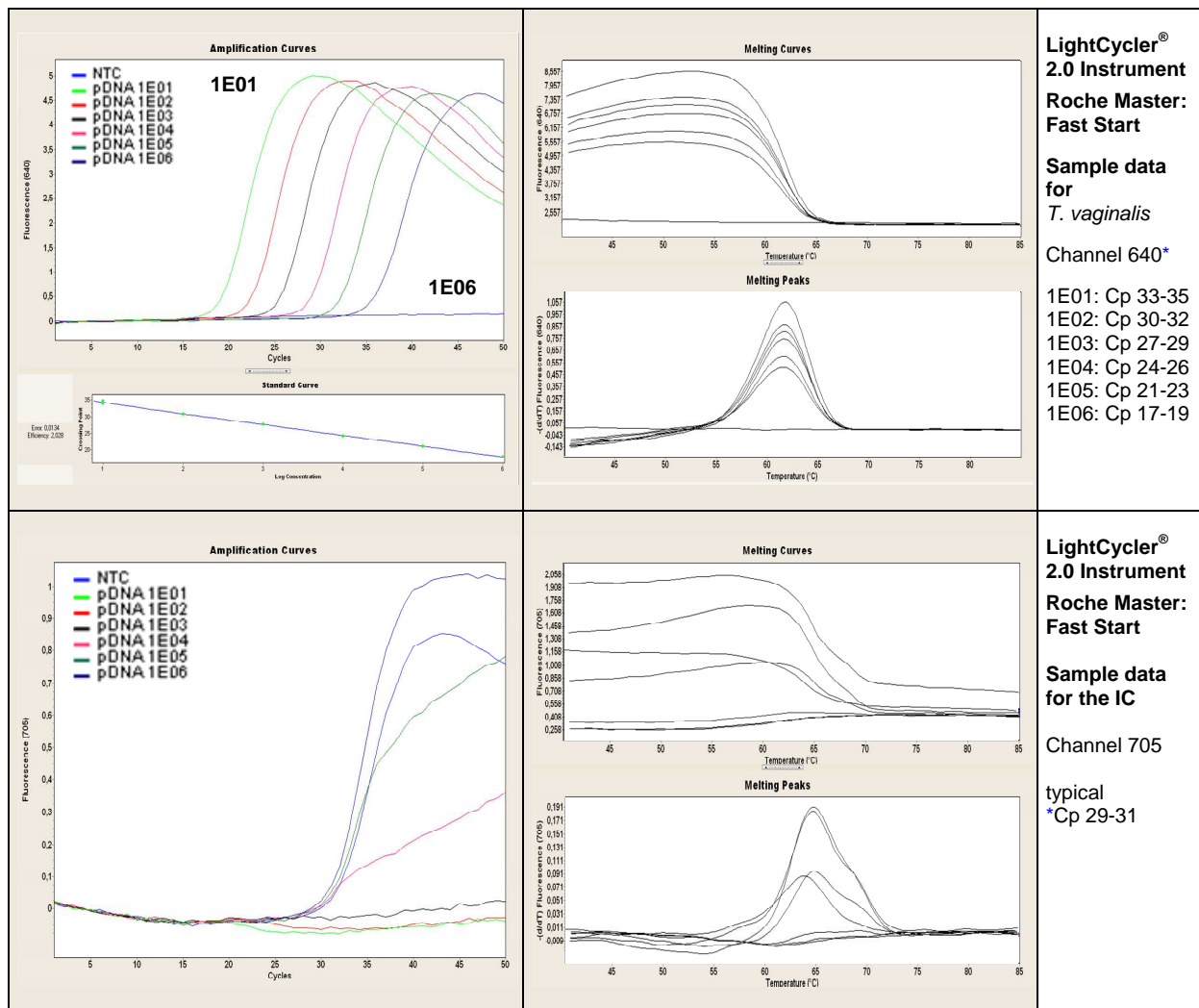


Fig.1. LightCycler® 2.0 sample data for the *T. vaginalis* detection system.

Upper panels: Left panel channel 640 quantification mode (Second Derivative Maximum) with amplification curves for *T. vaginalis*. Right panel channel 640 melting analysis for *T. vaginalis* (not relevant for detection).

Lower panels: Left panel channel 705 quantification mode (Second Derivative Maximum) for the IC. Right panel channel 705 melting analysis for the IC (not relevant for detection).

7.4. Interpretation of Data

Sample 640 <i>T. vaginalis</i>	Sample 705 IC	PositiveControl	Negative Control (NTC)	Result (warnings)
no amplification	detectable	amplification	negative	Negative (not detectable)
Cp < 37	not relevant	amplification	negative	Positive for <i>T. vaginalis</i>
no amplification	not detectable	amplification	not relevant	PCR failure , repeat experiment
not relevant	not relevant	no amplification	not relevant	PCR failure , repeat experiment
not relevant	not relevant	not relevant	positive	Contamination , repeat experiment

Table 3. Typical analysis results (LightCycler® 2.0 Instrument, Roche Master: Fast Start)

* **Note:** Fluorescence levels depend on instrument settings and may vary. The characteristics of the curve (shape and trend of fluorescence levels) should be similar to the curve in the manual. The fluorescent curves over cycles (quantification mode) must be smooth and not zigzag shaped. The Cp values will vary from instrument to instrument by up to 2 cycles, while the distance between two dilution steps should be relative constant (delta Cp). The Cp values described in this manual (chart text) have been obtained using the supplied standard row; the entire set of curves can be horizontal shifted.

8. LightCycler® 480 II / Cobas® Z480 Instruments

8.1. Programming

The protocol consists of four program steps

- 1: Denaturation: sample denaturation and enzyme activation
- 2: Cycling: PCR-amplification of the target DNA
- 3: Melting: melting curve analysis for identification of the PCR product derived from the target DNA
- 4: Cooling: cooling the instrument

Detection Format:

LightCycler® 480 II Instrument: 465-510, 498-640, 498-660

Cobas® Z480 Instrument: 465-510, 498-645, 498-700

Program Step:	Denaturation	Cycling			Melting			Cooling
Parameter								
Analysis Mode	None	Quantification mode			Melting Curves mode			None
Cycles	1	50			1			1
Target [°C]	95	95	62	72	95	40	85	40
Hold [hh:mm:ss]	00:10:00	00:00:05	00:00:05	00:00:15	00:00:30	00:02:00	00:00:00	00:00:30
Ramp Rate [°C/s] 96	4.4	4.4	2.2	4.4	4.4	1.5	-	1.5
Ramp Rate [°C/s] 384	4.6	4.6	2.4	4.6	4.6	2.0	-	2.0
Sec Target [°C]	-	-	55	-	-	-	-	-
Step Size [°C]	-	-	0.5	-	-	-	-	-
Step Delay (Cycles)	-	-	1	-	-	-	-	-
Acquisition Mode	None	None	Single	None	None	None	Continuous	None
Acquisitions [per °C]	-	-	-	-	-	-	1	-

Table 4

(Melting not relevant for detection)

8.2. Data Analysis

Note: Cobas® Z480 Instruments signal levels are about 50% compared to LightCycler® 480 II results.

Switch the color compensation mode on. If this mode is not enabled run the color compensation program. Follow the instructions in the manual of 'LightMix® Kit – Color Compensation 530/640/690.

Perform data analysis, as described in the LightCycler® Instrument operator's manual.

We recommend using the Second Derivative Maximum method (Automated (F" max). The cycle number of the Crossing Point (Cp) of each sample is calculated automatically. The Fit Points method is more-error prone due to the user's influence.

View *T. vaginalis* data with Filter Combination 498-640 Quantification mode. The negative control (NTC) must show no signal.

If the internal control is used, view data with Filter Combination 498-660, Quantification mode. The negative control and the low-concentrated *T. vaginalis* DNA samples (10 to 1.000 copies) should show an amplification curve for the IC with a Cp at approximately cycle 27-30.

The provided standard row of cloned and purified DNA with concentrations in the range from 10⁶ copies/rxn to 10¹ copies/rxn of *T. vaginalis* should have Cp values between cycles 17 and 36 (Cp values calculated with Second Derivative Maximum method).

8.3. Sample Data – Typical Results

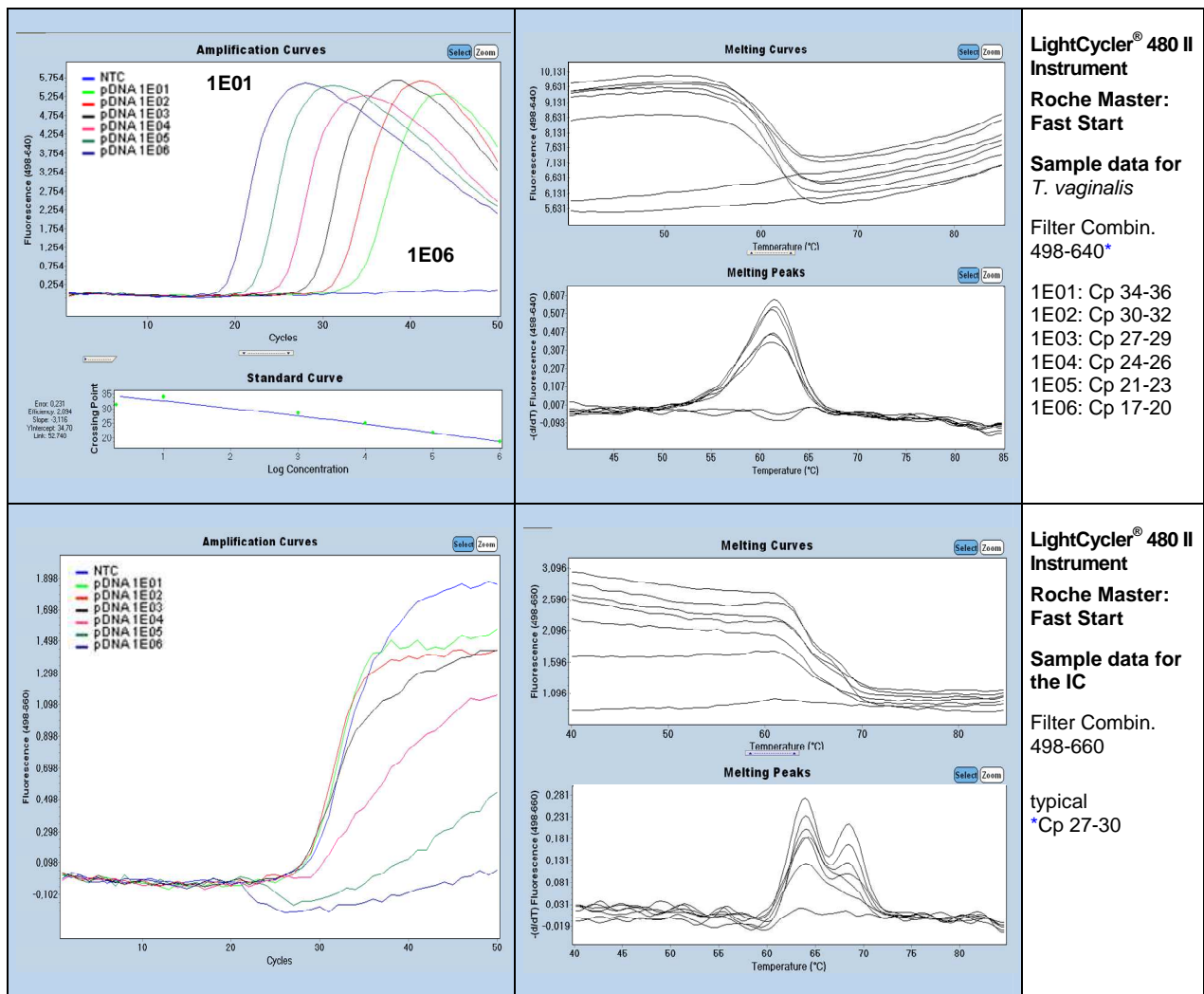


Fig.2. LightCycler® 480 II sample data for the *T. vaginalis* detection system.

Upper panels: Left panel Filter Combination 498-640 quantification mode (Second Derivative Maximum) with amplification curves for *T. vaginalis*. Right panel Filter Combination 498-640 melting analysis for *T. vaginalis* (not relevant for detection).

Lower panels: Left panel Filter Combination 498-660 quantification mode (Second Derivative Maximum) for the IC. Right panel Filter Combination 498-660 melting analysis for the IC (not relevant for detection).

* **Note:** Fluorescence levels depend on instrument settings and may vary. The characteristics of the curve (shape and trend of fluorescence levels) should be similar to the curve in the manual. The fluorescent curves over cycles (quantification mode) must be smooth and not zigzag shaped. The Cp values will vary from instrument to instrument by up to 2 cycles, while the distance between two dilution steps should be relative constant (delta Cp). The Cp values described in this manual (chart text) have been obtained using the supplied standard row; the entire set of curves can be horizontal shifted.

8.4. Interpretation of Data

Sample 640 <i>T. vaginalis</i>	Sample 660 IC	PositiveControl	Negative Control (NTC)	Result (warnings)
no amplification	detectable	amplification	negative	Negative (not detectable)
Cp < 38	not relevant	amplification	negative	Positive for <i>T. vaginalis</i>
no amplification	not detectable	amplification	not relevant	PCR failure, repeat experiment
not relevant	not relevant	no amplification	not relevant	PCR failure, repeat experiment
not relevant	not relevant	not relevant	positive	Contamination, repeat experiment

Table 5. Typical analysis results (LightCycler® 480 II Instrument, Roche Master: Fast Start)

9. Material Safety Data

According to OSHA 29CFR1910.1200, Commonwealth of Australia [NOHSC:1005, 1008 (1999)] and the European Union Directives 67/548/EC and 1999/45/EC any products which not contain more than 1% of a component classified as hazardous or classified as carcinogenic do not require a Material Safety Data Sheet (MSDS).

Product is not hazardous, not toxic, not IATA-restricted. Product is not from human, animal or plant origin. Product contains synthetic oligonucleotide primers and probes.

10. Version History

Notes in red mark events require to change procedures

V120214	Release version
V120523	Revised version
V121022	Correction in chapter 2 'Description' Roche Color Compensation reference removed.
V130813	Revised version

Roche SAP order n° 06896545001

Notice to Purchaser

A license under U.S. Patents 4,683,202, 4,683,195 and 4,965,188 or their foreign counterparts, owned by Hoffmann-La Roche Inc. and F. Hoffmann-La Roche Ltd ("Roche"), has an up-front fee component and a running-royalty component. The purchase price of this product includes limited, nontransferable rights under the running-royalty component to use only this amount of the product to practice the Polymerase Chain Reaction ("PCR") and related processes described in said patents solely for the research and development activities of the purchaser when this product is used in conjunction with a thermal cycler whose use is covered by the up-front fee component. Rights to the up-front fee component must be obtained by the end user in order to have a complete license. These rights under the upfront fee component may be purchased from Perkin-Elmer or obtained by purchasing an authorized thermal cycler. No right to perform or offer commercial services of any kind using PCR, including without limitation reporting the results of purchaser's activities for a fee or other commercial consideration, is hereby granted by implication or estoppel. Further information on purchasing licenses to practice the PCR process for research applications may be obtained by contacting the Director of Licensing at The Perkin-Elmer Corporation, 850 Lincoln Center Drive, Foster City, California 94404 or at Roche Molecular Systems, Inc., 1145 Atlantic Avenue, Alameda, California 94501. The purchase of this product does not convey any right for its use in clinical diagnostic applications. No rights for TaqMan technology under U.S. Patents 5,210,015 and 5,487,972 are hereby conveyed.

These reagents were developed and manufactured by TIB MOLBIOL GmbH, Berlin, Germany.
LightCycler® hybridization probes produced under license from Roche Diagnostics GmbH (worldwide excluding USA).

