

qPCR Master Mix DLP3 (2X)
Cat-No: S210 - 2x1,25 ml (100 rcs)

Features:

- The Master mix contains dUTP instead of dTTP
- The Mix contains ROX (500nM) as passive Reference dye (it provides a baseline in multiplex reactions)
- The qPCR / RTD-PCR Master mix DLP3 is ready-to-use and is optimized for high specificity and sensitivity because of optimized reaction buffer
- easy to us because ready-to-use Master Mix

Applications:

- Detection and quantification of DNA and cDNA targets
- Profiling gene expression
- Microbial detection
- Viral load determination

Description:

The Master Mix contains all reagents required for qPCR (except template and primer) in a premixed 2x concentrated ready-to-use solution. The high specificity and sensitivity of the mix is achieved by an optimized hot-start polymerase. Its activity is blocked at ambient temperature and switched on automatically at the onset of the initial denaturation. The thermal activation prevents the extension of non-specifically annealed primers and primer-dimer formations at low temperatures during PCR setup.

The mix offer dUTP instead of dTTP to prevent carry-over contaminations of DNA from previous PCR reactions.

Concentration: The Mastermix is 2x concentrated

List of components qPCR / RTD-PCR Master mix:

Hot-Start Polymerase for qPCR, dATP, dCTP, dGTP, dUTP, ROX, optimized reaction buffer with KCl and MgCl₂, stabilizers and enhancers, PCR-grade water

Transportation: with blue ice

Storage: at 4°C for 3 months, at -20°C for more than 12 months, **Note:** protect from Light

Usage:

| Components | Volume per reaction | final conc. |
|--|--|-------------|
| 2X qPCR / RTD-PCR Master mix DLP3 | 25 µl | 1x |
| Up-stream primer (10 µM stock) | 1,5 µl (range: 0,5-2.5 µl) | 300 nM |
| Down-stream primer (10µM stock) | 1,5 µl (range: 0.5-2,5 µl) | 300 nM |
| Template DNA | 5 µl (0.1-15 ng/ml plasmid DNA) (1-10 µg/ml genomic DNA) | < 500ng DNA |
| Sterile dest. Water (included) | up to 50 µl total reaction volume | |

- vortex all solutions carefully before using and before PCR
- may you add the enzyme mix after Template DNA
- an individual optimization of annealing temperature may be necessary for new combinations of primers and Template DNA

.. a good decision ..



General Thermo-Cycler protocol:

Note: working with EvaGreen just select the optical setting for FAM or SYBR Green at the cycler

| Step | Time | Temperature |
|--------------------------|--------------------|-------------|
| UNG treatment (optional) | 1x2 min | 50°C |
| Initial denaturation | 1-3 min | 95°C |
| 30-40 Cycles: | | |
| Denaturation | 15-30 sec | 95°C |
| Annealing | 30-65 sec | 55-65°C |
| Extension | 30 sec (per 500bp) | 72-75°C |

Note:

- an individual optimization of annealing temperature may be necessary for new combinations of primers and Template DNA

Ordering information.

| Cat.-no | Description | Amount |
|---------|----------------------|----------------|
| S210 | qPCR Master mix DLP3 | 100 rcs / 50µl |

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