

Product Information ExcelTaq™ series 2X Fast Q-PCR Master Mix (SYBR, ROX)

TQ1210 200 RXN

2X Fast Q-PCR Master Mix (SYBR, ROX) 1 ml x 2

TQ1211 500 RXN

2X Fast Q-PCR Master Mix (SYBR, ROX) 1 ml x 5

Storage

Aliquot to avoid multiple freeze-thaw cycles Protect from light

-20°C for 12 months

Features

- · High sensitivity and signal intensity
- Compatible with fast PCR program
- With smart blue contrast dye as a visual aid for reaction setup
- Low background
- · With ROX reference dve

Description

The ExcelTaq[™] 2X Fast Q-PCR Master Mix (SYBR, ROX) is a ready-to-use reagent with all the essential components for quantitative real-time PCR (qPCR) except primers and templates. The master mix features high sensitivity (Fig. 1) and signal intensity as well as low background and better compatibility with fast PCR program.

The ExcelTaq™ 2X Fast Q-PCR Master Mix (SYBR, ROX) contains hot-start *Taq* polymerase in an optimized buffer with dsDNA specific SYBR green fluorescent dye. This master mix allows sensitive, precise amplification, real-time tracking of the amplification process, and simultaneous quantification for targeted DNA molecules.

With inert smart blue contrast dye, the ExcelTaq™ 2X Fast Q-PCR Master Mix (SYBR, ROX) is ready-to-use and greatly reduces pipetting errors, while largely improving the reproducibility of the process. The ExcelTaq™ 2X Fast Q-PCR Master Mix (SYBR, ROX) also includes ROX reference dye for normalizing the fluorescent reporter signal in real-time quantitative PCR



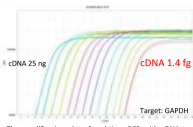


Fig. 1. The amplification plot of real-time PCR with cDNA template ranged from 25 ng to 1.4 fg in quantity, analyzed by using TQ1210 ExcelTaq™ 2X Fast Q-PCR Master Mix (SYBR, ROX) for qPCR amplification.

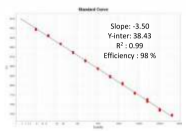


Fig. 2. The standard curve of TQ1210 ExcelTaq™ 2X Fast Q-PCR Master Mix (SYBR, ROX).

Application

- · Quantitative real-time PCR
- · Quantitative two-step real-time PCR

Instrument compatibility

- · Applied Biosystems system:
 - 7000, 7300, 7700,
 - 7900, 7900HT, 7900HT FAST
 - StepOne™ / StepOnePlus™

The Product is also compatible with several instruments that do not require the use of ROX as following.

- · BioRad system:
 - CFX96 / CFX384
 - Chromo 4[™] Real-Time Detector
 - Opticon[™] / Opticon[™] 2
- Cepheid system:
 - Smart Cycler®
 - Eppendorf system:
 - Mastercycler® ep realplex
- Roche system:
 - Roche LightCycler® 480 / Nano
- · QIAGEN system:
 - Rotor-Gene™ Q





Recommended primer design

· Amplicon size: 80-150 bp

• Tm value: around 60°C (calculated with Primer3 software)

• Primer length: 17-25 mer

• Sequence:

- 45-55% of GC content is recommended.
- Avoid regional high GC or AT content
- Avoid palindrome sequence
- Sequence with G or C at the 3' end is recommended.
- Specificity of primers should be confirmed through a BLAST search.

Recommended reaction mixture set up for qPCR

Template	2	2 μl*
Forward primer	50 – 400 n	M**
Reverse primer	50 – 400 n	M**
2X Fast Q-PCR Master Mix (SYBR,	ROX) 1	LO μl
H₂O	to 2	20 μΙ
Total volume	2	20 μΙ

^{*}Final template concentration varies depending on the copy number of target in the template solution. The recommended amount of template is: 100 fg -100 ng of cDNA, 80 pg -50 ng of gDNA or 10²-108

Recommended qPCR program

The Fast program is recommended for 2X Fast Q-PCR Master mix (SYBR, ROX). If the result is not good, please optimize the PCR program or try the standard program.

Fast program for qPCR

	' ' '			
	Steps	Temp.	Time	Cycles
	Template denature and enzyme activation	95°C	20 sec	1
	Denature	95°C	3 sec	40
	Annealing /Extension	60°C	30 sec	40
	Melting curve analysis	Refer to in	strument m	anual

Standard program for qPCR

Steps	Temp.	Time	Cycles
Template denature and enzyme activation	95°C	2 min	1
Denature	95°C	15 sec	40
Annealing/Extension	60°C	60 sec	40
Melting curve analysis	analysis Refer to instrument manual		

For more information, please refer to our website: www.smobio.com



molecules of plasmid.
**The PCR primer concentration for an optimal qPCR reaction may vary according to primers' properties and template condition.



Other Information

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Caution: Not intended for human or animal diagnostic or therapeutic uses.

Related Products

CK1000	Champion E. coli Transformation Kit
DM2300	ExcelBand 100 bp+3K DNA Ladder, 500 µl
DM3100	ExcelBand 1 KB (0.25-10 kb) DNA Ladder,
DIVISTOO	500 μl
DL5000	FluoroDye DNA Fluorescent Loading Dye
	(Green, 6×), 1 ml
NS1000	FluoroVue Nucleic Acid Gel Stain
	(10,000X), 500 μl
PM2510	ExcelBand Enhanced 3-color Regular
	Range Protein Marker, 250 μl × 2
RP1000	ExcelRT Reverse Transcriptase, 20,000 U
RP1100	ExcelRT One-step RT-PCR Kit, 50 RXN
RP1400	ExcelRT Reverse Transcription Kit II,
	100 RXN
RI1000	RNAok RNase Inhibitor, 2000 U
TF1000	SMO-HiFi DNA Polymerase, 100 U × 1
TP1000	ExcelTaq Taq DNA Polymerase, 500 U × 1
TP1200	ExcelTaq 5X PCR Master Dye Mix, 200 RXN
TQ1110	ExcelTaq 2X Q-PCR Master Mix (SYBR,
	ROX), 200 RXN
TQ2110	ExcelTaq 2X Q-PCR Master Mix (TaqMan,
	ROX), 200 RXN
WM1000	YesBlot Western Marker I, 250 μl

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