

PCRBIOSYSTEMS simplifying research

2x qPCRBIO SyGreen Mix with Fluorescein



Product description:

Combined with the latest advancements in polymerase technology and advanced buffer chemistry qPCRBIO SyGreen Mix offers market leading performance with minimal optimisation. qPCRBIO SyGreen Mix uses a proprietary intercalating dye which does not inhibit PCR, unlike other popular dyes.

qPCRBIO SyGreen Mix uses proprietary small molecular inhibitor technology that prevents formation of primer-dimers to improve reaction sensitivity and specificity.

High-throughput screening has resulted in a buffer system that allows efficient amplification from GC-rich and AT-rich templates, under fast and standard cycling conditions.

2x qPCRBIO SyGreen Mix with Fluorescein contains 20nM Fluorescein. This is for instruments which have a well-factor correction feature. Instruments which have this feature are Biorad iCycler®,MyiQ[™] and iQ®5 cyclers.

Pack Size	Format	Presentation
100 x 20µl rxns	2x ReadyMix	1 x 1ml
500 x 20µl rxns	2x ReadyMix	5 x 1ml
2000 x 20µl rxns	2x ReadyMix	20 x 1ml

Shipping and storage

On arrival the kit should be stored at -20°C. Avoid prolonged exposure to light. If stored correctly the kit will retain full activity for 12 months. The kit can be stored at 4°C for 1 month. The kit can go through 30 freeze/thaw cycles with no loss of activity.

Limitations of product use

The product may be used only for in vitro research purposes.

Technical support

For technical support and troubleshooting please email technical@pcrbio.com the following information:

Amplicon size Reaction setup Cycling conditions Screen grabs of amplification traces and melting profile

Instrument compatibility

Manufacturer	Instrument	Lo-ROX	Fluorescein
Bio-Rad®	iCycler®, MyiQ®, iQ ™5	Yes	Yes

Important considerations

Primer design: For efficient amplification under fast cycling conditions we recommend amplicon lengths between 80bp and 200bp. With all manufacturers master mixes the shorter the amplicon length the faster the reaction can be cycled. Amplicon lengths should not exceed 400bp. Primers should have a predicted melting temperature of around 60°C, using default Primer 3 settings (http://frodo.wi.mit.edu/primer3/).

Reaction setup

- 1. Before starting, briefly vortex 2x qPCRBIO SyGreen Mix.
- 2. Prepare a master mix based on following table:

Reagent	20µl reaction	Final concentration	Notes	
2x qPCRBIO SyGreen Mix	10µl	lx		
Forward primer (10µM)	0.8µl	400nM	See above for optimal primer design	
Reverse primer (10µM)	0.8µl	400nM		
Template DNA	<100ng cDNA, <1µg genomic	variable	See above for template considerations	
PCR grade dH,O	Up to 20µl final volume			

3. Program the instrument using following conditions, acquiring data on the SYBR® Green or FAM channel:

Cycles	Temperature	Time	Notes
1	95°C	2min	Polymerase activation, 2 minutes for cDNA and 3 minutes for genomic
40	95°C 60°C to 65°C	5 seconds 20-30 seconds	Denaturation Anneal/Extension, do not exceed 30 seconds, do not use temperatures below 60°C
Melt analysis	ysis Refer to instrument instructions		Optional melt profile analysis