

2x PCRBIO HS Taq Mix

Product description:

PCRBIO HS Taq Mix uses the latest developments in polymerase technology and buffer chemistry to enhance PCR speed, yield and specificity. The enzyme uses advanced hot-start technology for superior sensitivity. The enzyme and buffer system allow for superior PCR performance on complex templates such as mammalian genomic DNA. Due to enhanced efficiency and specificity the enzyme is perfectly suited to difficult PCR.

2x PCRBIO HS Taq Mix is a robust mix for all your everyday PCR applications including genotyping, multiplex PCR, screening, library construction, colony PCR and PCR direct from blood and urine. PCRBIO HS Taq DNA Polymerase can perform consistently well on a broad range of templates (including both GC and AT rich).

PCRBIO HS Taq Mix has an error rate of approximately 1 error per 2.0 x 10⁵ nucleotides incorporated. PCR products generated with PCRBIO HS Taq DNA Polymerase are A-tailed and may be cloned into TA cloning vectors.



2x PCRBIO HS Taq Mix 5x 1ml	25x 1ml

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.

PCRBIO HS Taq Mix is particularly resistant to PCR inhibitors. The mix is suitable for direct PCR from unprocessed samples including bacterial culture, bacterial colonies, blood and urine.

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

Important considerations

2x PCRBIO HS Taq Mix: The 2x mix contains PCRBIO HS Taq DNA Polymerase, 6mM MgCl₂, 2mM dNTPs, enhancers and stabilizers. It is not recommended to add further PCR enhancers or MgCl₂to the reaction. The buffer composition has been optimised to maximise PCR success rates.

Template: For eukaryotic DNA use between 5ng and 500ng per reaction, for cDNA use below 100ng per reaction.

Primers: Primers should have a predicted melting temperature of around 60°C, using default Primer 3 settings (http://frodo.wi.mit.edu/primer3/). The final primer concentration in the reaction should be between 0.2μ M and 0.6μ M.

Annealing: We recommend performing a temperature gradient to experimentally determine the optimal annealing temperature. Alternatively, we recommend a 55°C annealing temperature then increase in 2°C increments if non-specific products are present.

Extension: Optimal extension is achieved at 72°C. The optimal extension time is dependent on amplicon length and complexity of template. 15 seconds per kilobase (kb) is recommended for amplification from eukaryotic DNA for amplicons between 1kb and 6kb. For shorter amplicons a 1 second extension is sufficient.

Multiplex PCR: When first performing multiplex PCR it is recommended to run an annealing temperature gradient from 55°C to 65°C. The annealing temperature that results in the best specificity should be used in subsequent experiments. Fast cycling conditions should not be used for multiplex PCR. Initially, we recommend a 90 second extension time. This time may be further extended to increase yield.

Colony PCR: From bacterial colonies use a sterile tip to pick a colony and resuspend into a 50μ l reaction as described below. From liquid culture add 5μ l of overnight culture to the final mix. Increase initial denaturation time to 10 minutes.

Direct blood/urine PCR: Add 2µl mammalian blood or urine to a 50µl reaction as described below.

Reaction setup

1. Prepare a master mix based on the following table:

Reagent 50µl reaction		Final concentration	Notes	
2x PCRBIO HS Taq Mix	25.0µl	1x		
Forward primer (10µM)	2.0µl	400nM	See above for optimal primer design	
Reverse primer (10µM)	2.0µl	400nM		
Template DNA	<100ng cDNA, <500ng genomic variable		See above for template considerations	
PCR grade dH ₂ O	Up to 50µl final volume			

2. Cycle using conditions based on the following table:

Cycles	Temperature	Time	Notes
1	95°C	1min to 2min	Initial denaturation and enzyme activation. For colony PCR increase denaturation time to 10 minutes
40	95°C 55°C to 65°C 72°C	15 seconds 15 seconds 1 to 90 seconds	Denaturation Anneal Extension (15 seconds per kb). For multiplex PCR use 90 seconds