



REAGENTS FOR RESULTS

MyGo ProTaq Mix

Cat. No. 4152 | 7555

Component	80 Reactions (4152)	400 Reactions (7555)
2x MyGo ProTaq Mix	2 x 1ml	10 x 1ml

This product is for research use only

1. STORAGE

Store all components at -20°C with minimal exposure to light. If stored correctly the kit will retain full activity for 12 months. The kit may be stored at 4°C for short term use (1 month). The kit can go through up to 30 freeze/thaw cycles with no reduction in performance.

2. TECHNICAL ASSISTANCE

If you have any questions, or experience any difficulties with MyGo ProTaq Mix, please email reagentsupport@mygopcr.com, providing full details including amplicon size, reaction setup, cycling conditions and screen shots of gel images.

3. DESCRIPTION

MyGo ProTaq Mix is a high-performance mix designed for the amplification of difficult templates. The mix includes MyGo ProTaq Polymerase, dNTPs, and MgCl_2 in an optimised buffer designed to give the best results under challenging conditions such as the amplification of high GC targets and the presence of inhibitors.

MyGo ProTaq Polymerase uses antibody-mediated hot start to give fast, highly specific PCR. The enzyme is inactive at ambient temperatures, preventing the formation of primer-dimers and mis-primed products with the convenience of room temperature setup. The enzyme is activated at the start of a reaction with a 95°C incubation step. The enhanced efficiency and specificity of MyGo ProTaq Polymerase make it the ideal choice for the accurate amplification of long and complex templates.

MyGo ProTaq Mix is ready-to-use and supplied in a single tube containing all the reagents required for trouble-free PCR setup. It is recommended for the amplification of human genomic DNA up to 20kb. MyGo ProTaq Polymerase exhibits 2.5 fold higher fidelity than Taq DNA Polymerase. PCR products generated are A-tailed, and may be cloned into TA cloning vectors.



4. IMPORTANT NOTES

4.1 2x MyGo ProTaq Mix: The 2x mix contains MyGo ProTaq Polymerase, 6mM MgCl₂, 2mM dNTPs, enhancers and stabilizers. The buffer system has been optimised for the best PCR results and we do not recommend adding further enhancers or MgCl₂ to the reaction.

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

4.3 Primers: We recommend using primer design software Primer 3 (<http://frodo.wi.mit.edu/primer3/>). Primers should have a melting temperature (T_m) of approximately 60°C. The final primer concentration in the reaction should be between 0.2µM and 0.6µM.

4.4 Annealing: We recommend performing a temperature gradient to experimentally determine the optimal annealing temperature. As an alternative, you can use a 57°C annealing temperature then increase in 2°C increments if non-specific products are present.

4.5 Extension: The recommended extension temperature is 72°C. The optimal extension time will depend on amplicon length and complexity of template. For eukaryotic DNA we recommend 15 seconds per kilobase (kb) for amplicons below 5kb, and 40-60 seconds per kb for amplicons between 5-20kb.

REACTION SETUP

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

Component	50µl reaction	Final concentration	Notes
2x MyGo ProTaq Mix	25.0µl	1x	
Forward primer (10µM)	2.0µl	400nM	See 4.3 above
Reverse primer (10µM)	2.0µl	400nM	
Template DNA	<100ng cDNA <500ng gDNA	variable	See 4.2 above
PCR grade water	Up to 50µl total volume		

2. Cycle using conditions based on the following table:

Cycles	Temperature	Time	Notes
1	95°C	1 min to 2 min	Initial denaturation and enzyme activation
25-35	95°C	15 seconds	Denaturation
	55°C to 65°C	15 seconds	Anneal
	72°C	10 minutes*	Extension (50 seconds per kb). *See 4.5 above