AmpliTaq Gold[®] DNA Polymerase

S	Package Contents	Catalog Number N808-0240 AmpliTaq Gold® with Buffer I N808-0241 AmpliTaq Gold® with Buffer II 4311806 AmpliTaq Gold® with Gold Buffer Kit Contents	Size 250 Units 250 Units 250 Units
	Storage Conditions	 Store all contents at –20°C. 	
	Required Materials	 Template: cDNA, gDNA, λDNA 10 mM dNTP mix (Cat. no. 18427-088) Forward and reverse gene-specific prime: Autoclaved, distilled water E-Gel[®] General Purpose Gels, 1.2% (Cat. no. TrackIt[™] 1 Kb Plus DNA Ladder (Cat. no. 0.2 or 0.5-mL nuclease-free microcentrifuged 	no. G5018-01) 10488-085)
	Timing	Varies depending on amplicon length	
R	Selection Guide	PCR Enzymes and Master Mixes Go online to view related products.	
	Product Description	 AmpliTaq Gold[®] is derived from recombine 94 kDa DNA polymerase, encoded by a mathematical different buffers, which provide preferrent strength for PCR amplification reactions. With AmpliTaq Gold[®] DNA Polymerase, Face area provided into existing systems by modifying cycling parameters conditions for increased specificity, sensitifiered and a sensitive be completely or partially activated in a prostep, allowing flexibility and assembly of mathematical specificity. 	odified form of the It is available with d pH and ionic Hot Start and Time ng amplification or reaction vity, and yield. e state and can re-PCR heat
	Important Guidelines	 Take precautions to avoid cross-contamin aerosol-resistant barrier tips and analyzin a separate area from PCR assembly. If the samples contain EDTA or other che MgCl₂ concentration in the reaction mix p If proteases are present in the sample DN genomic DNA), inactivate the proteases h to 95°C for 5 minutes before adding Amp 	ag PCR products in lators, raise the proportionately. A (e.g. impure by heating samples
	Online Resources	Visit our product page for additional information and protocols. For support,	

visit www.lifetechnologies.com/support. For Research Use Only. Not for use in diagnostic procedures.



nzyme Characteristics

Hot-start:	Chemical
Length:	Up to 5 kb
Fidelity vs. Taq:	1X
Format:	Separate components

CR Reaction Setup

se the measurements below to prepare your PCR experiment, or enter your own arameters in the column provided.

Component	25-µL rxn	50-µL rxn	Custom		Final Conc.
Autoclaved, distilled water	to 25 μL	to 50 µL	to	μL	_
10X PCR Buffer I, II, or Gold Buffer	2.5 µL	5.0 µL		μL	1X
10 mM dNTP Mix*	0.5–5.0 μL	1.0 µL		μL	0.2 mM
25 mM MgCl ₂ **	1.5 µL	3.0 µL		μL	1.5 mM
10 µM forward primer	0.5 µL	1.0 µL		μL	0.2 µM
10 µM reverse primer	0.5 µL	1.0 µL		μL	0.2 µM
Template DNA	varies	varies			<1 µg/rxn
AmpliTaq Gold® DNA Polymerase (5 U/µL)	0.125 μL	0.25 μL		μL	1.25 U/ 50-μL rxn

Substituting dUTP for dTTP to control PCR product carryover may require higher oncentrations of dUTP (typically twice that of any other dNTP) for optimal amplification. Use MgCl, with Buffer II or Gold only. Buffer I already contains Mg.

CR Protocol

See page 2 to view a procedure for preparing and running your PCR experiment.

ptimization Strategies

Refer to the pop-up for guidelines to optimize your PCR reactions.

Limited Warranty, Disclaimer, and Licensing Information





AmpliTaq Gold[®] DNA Polymerase Protocol

The example PCR procedure below shows appropriate volumes for a single **50-µL** reaction. For multiple reactions, prepare a master mix of components common to all reactions to minimize pipetting error, and then dispense appropriate volumes into each 0.2–0.5 mL PCR reaction tube prior to adding template DNA and primers.

Timeline		Steps	Procedure Details					
1		Thaw reagents	Thaw, mix, and briefly centrifuge each component before use. Keep components on ice.					
		Prepare PCR master mix	Add the following components into appropriate wells or tubes.Note: Consider the volumes for all components listed in steps 2 and 3 to determine the correct amount of water required to reach your final reaction volume.Component50-µL rxnFinal Concentration Autoclaved, distilled water10X PCR Buffer I, II, or Gold Buffer5.0 µL1X					
2				10 mM dNTP mix		1.0 µL	0.2 mM each	
				25 mM MgCl, (with Buffer II or Gold only)			1.5 mM	
			Ŭ	AmpliTaq Gold [®] DNA Polymerase (5 U/µL)			1.25 U/rxn*	
			* The amount of AmpliTaq Gold [®] needed for the typical PCR amplification depends on cycling parameters. Start with 1.25 U/reaction.					
			Mix and briefly centrifuge the components.					
	300	Add template DNA and primers					Final Concentration	
				10 µM forward primer			0.2 μM	
3			· · · · · · · · · · · · · · · · · · ·	10 μM reverse primer Template DNA			0.2 μM < 1 μg/rxn*	
			Template DNAvaries< 1 µg/rxn** Preferably > 10 ⁴ copies of template but < 1 µg DNA/reaction.					
		Incubate reactions in a thermal cycler	Note: The pre-PCR heat step can completely activate AmpliTaq Gold [®] . If using the Time Release method, skip or reduce the duration of the initial denaturation step.					
			Step Tempera		ure (°C)	Time		
			Initial Denaturation		9		10 minutes	
4			25–35 PCR Cycles	Denature	95		15 seconds	
				Anneal	~55 (depending on primer T _m)		30 seconds	
				Extend	72		1 minute/kb	
			Final Ex	Final Extension 72		2	5 minutes	
			He	Hold 4			indefinitely	
5	And the second s	Analyze with gel electrophoresis	Analyze 10 µL using agarose gel electrophoresis. Use your PCR reaction immediately for down-stream applications, or store it at –20°C.					