



**SOLIS  
BIODYNE**

## FIREPol<sup>®</sup> DNA Polymerase Kit

Catalogue Number	Pack Size (5 U/ $\mu$ l)
01-01-KIT-0000S	100 U
01-01-KIT-00500	500 U
01-01-KIT-01000	1000 U
01-01-KIT-02000	2000 U



### Shipping:

At room temperature.

**Store at  $-20^{\circ}\text{C}$   
upon receipt**

### Batch Number and Expiry Date:

See vial.

### Storage and Stability\*:

- Routine storage at  $-20^{\circ}\text{C}$  ( $-28^{\circ}\text{C}$  to  $-18^{\circ}\text{C}$ ) until expiry date.
- Stable at  $4^{\circ}\text{C}$  ( $2^{\circ}\text{C}$  to  $8^{\circ}\text{C}$ ) for 6 months.
- Stable at room temperature ( $25^{\circ}\text{C}$ ) for 1 month.
- Freeze-thaw stability: 30 cycles.

### Reaction setup:

At room temperature.

Manufactured by Solis BioDyne, in compliance with the ISO 9001 and ISO 13485 certified Quality Management System.

## Product description:

- FIREPol<sup>®</sup> is a highly processive, thermostable DNA Polymerase. Due to its genetic modifications\*\*, it has enhanced stability at room temperature with no activity loss for up to 1 month. It is produced in *E. coli* strain that carries an overproducing plasmid with a modified gene of *Thermus aquaticus* DNA Polymerase.
- Possesses 5'→3' polymerase and 5'→3' exonuclease activity, as well as a non-template-dependent terminal transferase activity, but lacks a 3'→5' exonuclease (proofreading) activity making the generated product suitable for TA-cloning.
- Recommended for routine applications (generates fragments up to 3 kb from genomic DNA and 5 kb from low complexity samples, e.g., cDNA, plasmid). Is suitable for colony PCR.
- The fidelity of FIREPol<sup>®</sup> is similar to a regular *Taq* DNA Polymerase (error rate per nucleotide ca  $2.5 \times 10^{-5}$ ).

## Kit content:

Component	Catalogue Number			
	01-01-KIT-0000S	01-01-KIT-00500	01-01-KIT-01000	01-01-KIT-02000
FIREPol <sup>®</sup> DNA Polymerase (5 U/μl)	100 U / 20 μl	500 U / 100 μl	1000 U / 200 μl	2000 U / 400 μl
FIREPol <sup>®</sup> 10x Buffer B	500 μl	2.5 ml	5.0 ml	2 x 5.0 ml
FIREPol <sup>®</sup> 10x Buffer BD	500 μl	2.5 ml	5.0 ml	2 x 5.0 ml
25 mM MgCl <sub>2</sub>	500 μl	2.5 ml	5.0 ml	2 x 5.0 ml
10x GC-rich Enhancer	100 μl	100 μl	500 μl	500 μl

- FIREPol® DNA Polymerase (5 units/μl) in 20 mM Tris-HCl pH 8.7 at 25 °C, 100 mM KCl, 0.1 mM EDTA, 50% glycerol (v/v), and stabilizers.
- FIREPol® 10x Buffer B (Mg<sup>2+</sup> free, with detergent): 0.8 M Tris-HCl, 0.2 M (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 0.2% w/v Tween-20.

FIREPol® 10x Buffer B contains non-ionic detergent suppressing inhibitory effects of the traces of DNA extraction buffer and enhancing PCR yield and efficiency.

- FIREPol® 10x Buffer BD (Mg<sup>2+</sup> and detergent free): 0.8 M Tris-HCl, 0.2 M (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>.
- 25 mM MgCl<sub>2</sub>
- 10x GC-rich Enhancer is an additive that facilitates amplification of difficult templates (e.g. GC-rich DNA templates).

This solution should be used at a defined final concentration (1x, 2x or 3x solution). 10x GC-rich Enhancer is NOT a reaction buffer and should be used ONLY IF non-specific amplification occurs.

### Additional reagents required:

- Template DNA
- Gene-specific primer pair
- dNTP Mix (20 mM of each, Cat. No. 02-31-00020)
- Nuclease-free PCR Grade Water (Cat. No. water-025)

## Step-by-step guidelines:

1. Thaw the reagents at room temperature. Mix each reagent by gentle vortexing or pipetting up and down, then spin down.
2. Prepare a reaction mix at room temperature. Add all required components except the template DNA.

Component	Volume <sup>1</sup>	Final conc.
FIREPol <sup>®</sup> DNA Polymerase (5 U/ $\mu$ l)	0.08–0.2 $\mu$ l	0.02–0.05 U/ $\mu$ l
FIREPol <sup>®</sup> 10x Buffer B or BD	2 $\mu$ l	1x
25 mM MgCl <sub>2</sub>	1.2–2 $\mu$ l	1.5–2.5 mM
dNTP Mix (20 mM of each)	0.2 $\mu$ l	200 $\mu$ M of each
Forward Primer (10 $\mu$ M)	0.2–0.6 $\mu$ l	100–300 nM
Reverse Primer (10 $\mu$ M)	0.2–0.6 $\mu$ l	100–300 nM
10x GC-rich Enhancer (optional)	2, 4 or 6 $\mu$ l	1x, 2x or 3x
Template DNA (added at step 4)	Variable	Variable <sup>2</sup>
Nuclease-free water	up to 20 $\mu$ l	
<b>Total reaction volume</b>	<b>20 <math>\mu</math>l</b>	

<sup>1</sup> Scale all components proportionally according to sample number and reaction volumes. Make sure you use enough of each reagent for your reactions, plus 10% extra volume to accommodate pipetting errors.

<sup>2</sup> For low complexity templates (i.e. plasmid, lambda), use 20 pg–2 ng of DNA per 20  $\mu$ l reaction. For higher complexity templates (i.e. gDNA), use 2 ng–200 ng of DNA per 20  $\mu$ l reaction.

3. Mix the reaction mix thoroughly, then spin down. Dispense appropriate volumes of mix into PCR wells or tubes.
4. Add template DNA to the PCR wells. Seal the wells using the procedure recommended for the cycling instrument being used and centrifuge the reactions briefly.
5. Incubate your PCR reactions in thermal cycler as follows.

Step	Temperature	Time	Cycles
Initial denaturation <sup>1</sup>	95°C	3–5 min	1
Denaturation	95°C	15–30 sec	26–35
Annealing <sup>2</sup>	50–68°C	30–60 sec	
Extension <sup>3</sup>	72°C	45 sec–4 min	
Final extension	72°C	5–10 min	1

<sup>1</sup> Complex templates, such as gDNA, require longer time to denature (5 min). With low complexity templates (i.e. lambda, plasmid DNA), initial denaturation time can be reduced to 3 min.

<sup>2</sup> The annealing temperature depends on the melting temperature of the primers.

<sup>3</sup> Extension time depends on the length of the fragment to be amplified. A time of 1 min/kb is recommended.

## Recommendations for a successful PCR experiment

Prerequisites for a successful PCR include the design of optimal primers, the use of high-quality template DNA and appropriate concentrations of reaction components.

Use dedicated software, such as Primer3 and NCBI Primer-BLAST to design target-specific primers. The optimal primer length is 20–30 bp, with GC-content 35–65% and calculated melting temperatures ( $T_m$ ) 60–70°C.  $T_m$  of the two primers should not differ by more than 3°C. Analyze your primers for self-

complementarity and stable secondary structures, presence of secondary structures increases probability of mis-priming and primer-dimers formation.

The integrity, purity and concentration of the DNA template should be suitable for the PCR experiment. Always include a no-template control (NTC) by replacing the DNA template with the same volume of nuclease-free water.

## Troubleshooting Guide

### No or low PCR yield

- Cycling conditions are not optimal – adjust the primer annealing temperature ( $T_a$ ); if needed determine the optimal  $T_a$  by running a temperature gradient; increase the extension time (if amplifying a long target); increase the number of cycles by 3–5.
- Poor quality of template – check the template's purity and integrity, ensure that your template doesn't contain PCR inhibitors.
- Template concentration is too low – increase the concentration of DNA template in a reaction by increasing sample volume; or concentrate the DNA by precipitation before adding the sample to the reaction.
- Primer concentration is not optimal – titrate primer concentration (final concentration 100–300 nM of each); ensure that both primers have the same concentration.
- Reaction components are degraded – check the storage conditions and expiry date of the reagents; perform a positive control with template DNA and/or reagents previously known to amplify.

## Non-specific products

- Non-specific amplification – use hot-start PCR enzyme (e.g., HOT FIREPol® DNA Polymerase, Cat. No. 01-02-00500); ensure that your primers are target specific.
- Primer concentration is not optimal – titrate primer concentration (final concentration 100–300 nM of each); too high primer concentration can reduce the binding specificity, resulting in unwanted products.
- Primer annealing temperature ( $T_a$ ) is too low – increase the  $T_a$ ; keep your primer annealing temperature 2–5°C below the  $T_m$  of the primer having the lowest  $T_m$ .
- Too many cycles – reduce the cycle number by 3–5.
- Contamination – to avoid contamination, work in dedicated space, keep pre- and post-amplification areas separate, decontaminate your surfaces and equipment, if possible, aliquot your reagents into smaller volumes to prevent contamination of stock solutions.

## Smearing in electrophoresis

- Too much template – load lower amount or prepare serial dilutions of template.
- Too many cycles – reduce the cycle number by 3–5.
- Extension time is too long – reduce extension time.
- Primer design is not optimal – review your primers and redesign the primers if needed.
- Enzyme concentration is too high – decrease the amount of enzyme in final solution by 0.005 U/ $\mu$ l increments (optimal polymerase concentration in final PCR solution is 0.02–0.05 U/ $\mu$ l).



## Unit definition:

One unit is defined as the amount of enzyme required to catalyze the incorporation of 10 nmol of dNTPs into an acid-insoluble form in 30 minutes at 74°C.

## Safety precautions:

Please refer to the Safety Data Sheet for more information.

## Technical support:

Contact your sales representative for any questions or send an email to [support@solisbiodyne.com](mailto:support@solisbiodyne.com)

**DS-01-01-KIT v2. Effective from: 02.04.2024**

**Reason for revision:** Kit component 10x Solution S changed to 10x GC-rich Enhancer.

\*Product stability is assessed using routine QC assays and QC criteria set forth in the product specification and are intended to provide guidelines for shipping and storage conditions only. The customer or its designee shall be responsible for conducting all necessary stability testing applicable to their assay and/or QC criteria, and to comply with any applicable regulatory requirements or guidelines. Such stability testing shall include testing to validate the lead times for shipment, the shelf life of, and the product specifications applicable to shipment, storage and handling of the assay assembled and packed by the customer.

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**Trademark information:** FIREPol is a registered trademark of Solis BioDyne OÜ.

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