



## Klentaq-S      Cat #: 105

**Amount:** 100 µl enzyme (sufficient for 2000 x 25 µl reactions up to 1 kb)

**Shipping conditions:** Ambient

**Storage conditions:** -20°C

### PRODUCT DESCRIPTION:

Klentaq-S is a mutant of Klentaq that has the feature of incorporating both dNTPs and ddNTPS. It can be used in Pyrophosphorolysis-Activated Polymerization (PAP) for excellent specificity of primer binding.

10x buffer composition is: 500 mM Tris-Cl pH 7.8, 160 mM ammonium sulfate, 0.25% Brij 58, and 35 mM magnesium chloride. The enzyme may not perform as well at a higher pH.

### TYPICAL PROTOCOL for Pyrophosphorolysis-Activated Polymerization (PAP) for a 25 µl reaction:

Reagent	Volume	Final Concentration
10x Klentaq-S reaction buffer	2.5 µl	1x
dNTP mix (10 mM)	0.0625 - 0.5 ul	25-200 uM each
Left Primer	variable	25-200 nM
Right Primer	variable	25-200 nM
Na <sub>4</sub> PPi	variable	90 uM
DMSO	variable	2%
BSA (optional)	variable	0.15 mg / ml
DNA template†	variable	100-200 ng
Klentaq-S**	0.05 – 0.25 µl **	
De-ionized distilled H <sub>2</sub> O	Adjust final volume to 25 ul	-

† DNA amount depends mostly on genome size and target gene copy number.

\*\* To determine specific optimal enzyme concentration, we strongly recommend an enzyme titration test for each target. Targets larger than 1 kb may require more enzyme or may benefit from the LA (Long-Accurate) version of the polymerase.

### CYCLING CONDITIONS\*

Initial denaturing:      95° for 2 minutes

25 “Touchdown” cycles: 94° for 15 seconds  
                                   60° for 30 seconds  
                                   64° for 30 seconds  
                                   68° for 1 minute  
                                   72° for 1 minute

\*Suggested conditions for PAP for 25 ul reactions. Optimal temperatures may vary depending on primer sequence. Extension times may be increased for longer targets. We typically recommend 1 minute + 1 minute per kb target.

**Please visit us on the web at [www.klentaq.com](http://www.klentaq.com) for troubleshooting and detailed protocols.**

### References:

Liu Q and Sommer SS. (2002) Pyrophosphorolysis-activatable oligonucleotides may facilitate detection of rare alleles, mutation scanning and analysis of chromatin structures. *Nucleic Acids Res.* 30(2):598-604.

Liu Q, et al. (2006) Multiplex dosage pyrophosphorolysis-activated polymerization: application to the detection of heterozygous deletions. *Biotechniques* 40(5):661-8.