Hot Start CesiumTaq

Amount: 100 μl (2000 x 25 μl reactions) **Shipping conditions:** Ambient temperature

Storage conditions: -20°C

Thermostability: Retains at least 85% activity after 1 hour at 95°C

Expiration: On tube label



PRODUCT DESCRIPTION:

Hot Start CesiumTaq is made with aptamer-based technology, enabling room temperature reaction set-up.

CesiumTaq is a double cold-sensitive mutant of Taq DNA polymerase. Due to its suppressed activity at low temperatures this enzyme is designed for hot-start PCR performance. Addition of the aptamer further enhances its hot start capabilities. The aptamer binds to the polymerase at sub-cycling temperatures, inactivating the enzyme and preventing spurious amplification.

Cat #: HS200

10x buffer composition is: 500 mM Tris-Cl pH 8.3, 160 mM ammonium sulfate, 0.25% Brij 58, and 25 mM magnesium chloride.

TYPICAL PCR PROTOCOL for a 25 µl reaction:

Reagent	Volume	Final Concentration
10x Taq Mutant Reaction Buffer	2.5 μl	1x
dNTP mix (10 mM each)	0.5 μ1	200 μM each
Left Primer	variable	200 nM
Right Primer	variable	200 nM
DNA template†	variable	0.1-100 ng
PCR Enhancer Cocktail (recommended)*	12.5 μl	1x
Hot Start CesiumTaq	0.05 - 0.25**	1 unit
De-ionized distilled H2O	Adjust final volume to 25 μl	

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

CYCLING CONDITIONS:

1. Denaturing: 94° for 2-8 minutes for 1 cycle

2. Denaturing: 94° for 40-60 seconds

3. Annealing: 50°-68° depending on the specific Tm primers for 40-60 seconds

4. Extension: 68° for 2 min/kb target

5. Repeat steps 2-4 for 25-40 cycles

Please visit us on the web at www.klentaq.com for troubleshooting and detailed protocols.

REFERENCES:

Kermekchiev, M.B., et al. (2003) Cold-sensitive mutants of Taq DNA polymerase provide a hot start for PCR. Nucl Acids Res. 31, 6139-6147.

^{*} For optimal performance, we recommend using one of our PCR Enhancer Cocktails (PEC-a, PEC1-GC, PEC-2, or PEC-2-GC) which are specially formulated for use with whole blood, serum, or plasma or 1.3M Betaine, a general PCR enhancer.

^{**} To determine specific optimal enzyme concentration, we strongly recommend an enzyme titration test for each target A good starting amount of the enzyme per 25 μ l reaction is 0.05 μ l for purified DNA templates and 0.25 μ l for crude samples containing 5-10% whole blood, plasma or serum. Targets larger than 1 kb may require more enzyme or may benefit from the use of an LA (Long Accurate) version of the polymerase.