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NORTH AMERICA

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Diamond *Taq*[®] DNA Polymerase Specification Sheet Reference: TAQ-I021

Eurogentec products are sold for research or laboratory use only and are not to be administrated to humans or used for medical diagnostics.

For medical diagnostics, please use the TAQ-I020 references.

Source

Diamond *Taq*[®] is a highly thermostable enzyme produced and purified from recombinant *Escherichia coli* bacterium containing the *Thermus aquaticus* DNA Polymerase gene.

Intended use

Diamond *Taq*[®] is particularly suited for PCR applications that require high sensitivity and ultra low level of bacterial & fungal DNA. The GMP manufacturing & purification processes minimize the risk of false positive results due to residual DNA contamination (bacterial or fungal). The enzyme is QC-tested to verify that < 1fg of genomic *E. coli* DNA (or 0.2 copy) is present in a standard aliquot containing 1 unit of *Taq*. Bioburden is guaranteed \leq 10 CFU/ml, but is typically = 0 CFU/ml.

Package contents

Reference	Units	Volume	Concentration	Volume Diamond <i>Taq</i> [®] reaction buffer (10 X)*	Volume 25 mM MgCl ₂
TAQ-I021-100 (sample)	100	20 μ l	5 U/ μ l	1 ml	1 ml
TAQ-I021-1000	1000	200 μ l	5 U/ μ l	6 ml	6 ml
TAQ-I021-5000	5000	1 ml	5 U/ μ l	30 ml	30 ml
TAQ-I021-25000	5 x 5000	5 x 1 ml	5 U/ μ l	5 x 30 ml	5 x 30 ml

*750 mM Tris-HCl pH 8.8 (at 19°C), 200 mM (NH₄)₂SO₄, 0.1% (v/v) Tween 20 and stabilizer.

Storage conditions

Storage at -20°C is recommended

Storage and dilution buffer

20 mM Tris-HCl, 1 mM DTT, 0.1 mM EDTA, 0.1 M KCl, 0.5% (v/v) Nonidet P40, 0.5% (v/v) Tween 20, 50% (v/v) glycerol, pH 8.0 (19°C) and stabilizer.

Enzyme Specifications

Each lot of enzyme, buffer and MgCl₂ is functionally tested and quality controlled to ensure the following specifications of the IVD-GMP products.

Appearance	Clear, colourless solution
Identity (SDS-PAGE)	MW approx. 95 kDa
Volume activity	\geq 5 U/ μ l
Purity (SDS-PAGE)	> 98%
Performance test: PCR on λ DNA	0.5 kb fragment positive down to 5 pg
Performance test: PCR on genomic DNA	0.1 kb fragment positive down to 10 pg
Ribonucleases (up to 10 U, 1h, 37 °C)	Not detectable
Endonucleases (up to 30 U, 16h, 65 °C)	Not detectable
Exonucleases (up to 30 U, 16h, 65 °C)	Not detectable
Nicking activity (up to 30 U, 16h, 65 °C)	Not detectable
Nicking activity (up to 30 U, 16h, 65 °C)	Not detectable
<i>E. coli</i> residual DNA	< 1 fg / Taq Unit
Bioburden	\leq 10 CFU/ml
Stability	24 months (at-20°C) from date of manufacture
Animal-derived additives	None

Unit definition

One unit is defined as the amount of enzyme that incorporates, 10 nmoles of dNTPs into acid insoluble form in 30 minutes at 74 °C.

Shipping conditions

Shipping at room temperature

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Reaction Conditions

For a 100 µl Reaction

Diamond Taq [®] Reaction Buffer (10x)	10 µl
MgCl ₂ solution	6 µl (1.5 mM)
Diamond Taq [®]	0.8 to 2.5 units
dNTP	200 µM each dNTP
Primers	0.1 nmol each
H ₂ O	As required
DNA template	As required

Magnesium

This DNA polymerase is a magnesium-dependent enzyme. We recommend increasing the magnesium concentration for long DNA fragments. Excess Mg²⁺ stabilizes the DNA double strand and consequently prevents complete denaturation of DNA, which reduces the extension yield. It may also stabilize spurious primer/template annealing, thus decreasing specificity.

Recommendation

Homogenize Diamond Taq[®] solution by flipping the tube 4 to 5 times.

Cycling conditions

Classical PCR protocol used for 500 bp lambda DNA amplification*

25 cycles	{	95°C	3 min
		94°C	30 sec
		T _m -2°C	30 sec
		72°C	1 min/kb
		72°C	7 min
		4°C	end temperature

**Condition will vary from reaction to reaction and may need optimization for maximal performances. Duration and temperature for denaturation and annealing steps depend on the type of cyler and primers design. We advise you to check primer design by using primer design software.*

Disclaimer

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