

# FastGene® apTaq HotStart Polymerase

Technical Data Sheet

#### **Product Description**

The FastGene® apTaq HotStart Polymerase uses an aptamer based HotStart activation technology. The aptamer based HotStart inhibits any polymerase acitivity at temperatures below 45 °C and has an immediate activation.

### **Product Applications**

The FastGene® apTaq HotStart Polymerase is ideally suited for:

- Fast PCR
- Routine PCR
- · PCR using complex templates
- SNP Analysis
- Any standard PCR application for which a hot start formulation of a high-quality thermostable DNA polymerase is required.

# Shipping and Storage

FastGene® apTaq HotStart Polymerase kits are shipped on ice packs. Upon arrival, store kit components at -20 °C, in a constant-temperature freezer. When stored under these conditions and handled correctly, full activity of the kit is retained until the expiry date indicated on the kit label.

#### Handling

Always ensure that the product has been fully thawed and mixed before use. Reagents may be stored at 4 °C for short-term use (up to 1 month). Return to -20 °C for long-term storage. Please make sure to centrifuge briefly each tube prior usage.

Kit Codes and Components						
LS34s LS34	FastGene® apTaq HotStart Polymerase FastGene® apTaq HotStart Polymerase	Sample 500 Units				
Related Products						
L\$20 L\$21 L\$22 L\$23 L\$24 L\$25 L\$26 L\$27 L\$28 L\$29 L\$30 L\$31	FastGene® Taq DNA Polymerase FastGene® Taq DNA Polymerase FastGene® Taq DNA Polymerase FastGene® HotStart TAQ DNA Polymerase FastGene® TAQ Ready Mix PCR Kit FastGene® TAQ Ready Mix PCR Kit FastGene® Optima FastGene® Optima FastGene® Optima FastGene® Optima FastGene® Optima FastGene® Optima	100 Units 500 Units 2000 Units 100 Units 250 Units 1000 Units 50 x 50µl rxns 250 x 50µl rxns 250 Units 500 x 25 µl rxns 50 Units 20 x 25 µl rxns				
Direct PCR						
LS05 LS06	DNAreleasy Advance DNAreleasy Advance	10 preps 50 preps				
Quick Notes						

#### Quick Notes

- · Replace Taq for Standard PCR, with no protocol change.
- Buffer is supplied at 5x concentration with separate magnesium for optimization.
- Products can be cloned in T-overhang cloning vectors.

#### Optimizing MgCl,

+49 2421 554960

Aptamer-based HotStart polymerase and/or reactions with increased amounts of template (e.g., genomic DNA), primer and nucleotides might require higher MgCl<sub>2</sub> concentrations. A separate 50 mM MgCl<sub>2</sub> solution is supplied with the enzyme. Please adjust the MgCl<sub>2</sub> concentration according to the table below:

Final conc. in 50 µl reaction	2 mM	2.5 mM	3 mM	3.5 mM	4 Mm
Volume of 50mM MgCl <sub>2</sub> Stock	2 µl	2.5 μΙ	3 μΙ	3.5 µl	4 μl



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# FastGene® apTaq Protocol

FastGene® apTaq HotStart Polymerase can be used to replace any commercial Taq DNA polymerase in an existing protocol. To allow the most seamless integration of FastGene® apTaq HotStart Polymerase into existing protocols, be sure to match reaction conditions, particularly the MgCl<sub>2</sub>, primer and enzyme concentrations, as closely as possible.

### **Step 1: Prepare the PCR master mix**

- Ensure that all reagents are properly thawed and mixed.
- Prepare a PCR master mix containing the appropriate volume of all reaction components common to all or a subset of the reactions to be performed.
- Calculate the required volumes of each component based on the following table:

Component	50 µl rxn¹	Final conc.
PCR-grade water	Up to 50 µl	N/A
5 U/µl of FastGene® apTaq	0.5 μl²	2.5 U <sup>2</sup>
5X FastGene® apTaq Buffer	10 µl	1X
50 mM MgCl <sub>2</sub>	2 μl³	2 mM³
dNTP Mix (10 mM each)	1 µl	0.2 mM each
Forward Primer (10 µM)	2.5 µl	0.5 µM
Reverse Primer (10 µM)	2.5 µl	0.5 μΜ
Template DNA <sup>4</sup>	As required	As required

 $<sup>^{1}</sup>$  Reaction volumes of  $10-50~\mu l$  are recommended. For volumes smaller than  $50~\mu l$ , scale reagents down proportionally.

# Step 2: Set up individual reactions

- Transfer the appropriate volume of PCR master mix, template and primer to individual PCR tubes/wells or a PCR plate.
- Cap or seal individual reactions, mix and centrifuge briefly.

Step 3: Run the PCR

Step	Temperature	Duration	Cycles
Initial denaturation	95 °C	1 min¹	1
Denaturation	95 °C	30 sec	
Annealing <sup>2</sup>	T <sub>m</sub> + 2 °C	15-30 sec	20-40³
Extension	72 °C	1 min/kb	
Final extension (optional) <sup>4</sup>	72 °C	1 min/kb	1
Store	4 – 10 °C	HOLD	1

 $<sup>^1</sup>$  Initial denaturation for 1 min at 95  $^{\rm oC}$  is recommended for most assays. For GC-rich targets (>65% GC), 5 min at 95  $^{\rm oC}$  may be used.

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For technical support please contact:

info@nippongenetics.eu

 $<sup>^2</sup>$  For small to midrange products, half the concentration of enzyme can be used. For GC-rich and other difficult templates, higher enzyme concentrations (up to 5 U per 50  $\mu l$  reaction) may be required.

<sup>&</sup>lt;sup>3</sup> A final MgCl<sub>2</sub> concentration of 2 mM is sufficient for most standard applications. For assays that do not perform well with 2 mM MgCl<sub>2</sub>, the optimal MgCl<sub>2</sub> concentration for each primer–template combination should be determined empirically.

<sup>&</sup>lt;sup>4</sup> ≤100 ng for genomic DNA; ≤1 ng for less complex DNA (e.g. plasmid, lambda).

 $<sup>^2</sup>$  An annealing temperature 2  $^{\circ}$ C higher than the calculated melting temperature ( $T_m$ ) of the primer set is recommended as a first approach. If low yields and/or nonspecific amplification is obtained, an annealing temperature gradient PCR is recommended to determine the optimal annealing temperature of the primer pair (gradient should be at least between  $-2^{\circ}$ C and  $+3^{\circ}$ C).

<sup>&</sup>lt;sup>3</sup> 35 cycles are sufficient for most assays. A higher number of cycles may be necessary for assays requiring higher sensitivity, while lower cycle numbers can be used if the template copy number is high.

<sup>&</sup>lt;sup>4</sup>Final extension should be included if PCR products are to be cloned into TA cloning vectors.