

Nova Taq Premix
2 fold concentrated
Reference: AB12006; AB12007; AB12008



INTENDED USE AND PRESENTATION:

This Taq Premix is widely used in molecular biology.

AB12006, 1 mL. 1mL.

AB12007, 3 mL. 1mL x 3.

AB12008, 5 mL. 1mL x 5.

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

For research use only.

SUMMARY, EXPLANATION AND LIMITATIONS:

Nova Taq Premix is a premixed solution containing everything needed for successful PCR reaction except specific primers and DNA template.

The mix includes high-quality recombinant Nova Taq DNA polymerase, nucleotides and magnesium in a PCR reaction buffer.

For the reaction set-up add the PCR Mix (10 or 25 μ L) to the primers, template and water for the total reaction volume of 20 or 50 μ L.

Quality Control: Activity and stability tested at 20, 30 and 40 cycles of PCR reactions at 95°C. Free of detectable, non-specific nucleases.

Several cycles of freezing/thawing are allowed.

APPLICATIONS:

Nova Taq Premix is a premix for all your everyday PCR reactions, gene expression profiling, Microbial & Viral pathogen detection. This Nova Premix allows efficient amplification from GC-rich and AT-rich templates, under fast and standard cycling conditions.

PRODUCT COMPOSITION:

-Nova Taq DNA Polymerase 1 unit/ μ L in reaction buffer, 4 mM MgCl₂, 0,4 mM of each dATP, dCTP, dGTP, dTTP.

-Nuclease free water.

METHODS AND PROCEDURE:

Optimal reaction conditions, such as reaction time, temperature, and amount of template DNA, may vary and must be individually determined.

General Reaction Protocol:

1. Thaw 2X Nova Taq Premix solution.
2. Prepare a master mix.

Component	Volume	Final Conc.
2X Nova Taq Premix	10 μ L	1X
Upstream Primer (10 pmoles/ μ L)	0,3~1,0 μ L	0,1~1,0 μ M
Downstream Primer (10 pmoles/ μ L)	0,3~1,0 μ L	0,1~1,0 μ M
Template DNA	Variable	1 ~100 ng/ μ L
Sterilized D.W.	Up to 20 μ L	-
Total Volume	20 μ L	-

*Primers should have a predicted melting temperature of around 60°C, using default Primer 3 settings (<http://frodo.wi.mit.edu/primer3/>).

Amount of template:

Bacteriophage λ , cosmid, plasmid \rightarrow 10 fg~300 ng

Total genomic DNA \rightarrow 100 ng~1 μ g

3. Mix the master mix thoroughly and dispense appropriate volumes into PCR tubes. Centrifuge the reactions in a microcentrifuge for 10 seconds.

4. Perform PCR using your standard parameters (3-step cycling).

Step	Temperature & Reaction Time		
Initial denaturation	3~5 min. at 95°C	-	-
25~35 cycles	30~60 sec. at 95°C	30~60 sec. at 50~68°C	1~4* min. at 72°C
Final extension	-	-	5~10 min at 72°C

*For PCR products longer than 3~4 Kb, use an extension time of approximately 1 min per Kb DNA.

5. Separate the PCR products by agarose gel electrophoresis and visualize with EtBr or any other means.

Los fragmentos de ADN amplificados con Nova Taq ADN Polimerasa presentan terminaciones en A, permitiéndole hacer clones usando el vector-T.

REQUIRED MATERIALS BUT NOT SUPPLIED:

All reagents, materials, and laboratory equipment for qPCR procedures are not provided with this polymerase. This includes sterile reaction tubes, micropipettes and tips, template DNA, gen-specific PCR primer pair, dNTPs mixture, PCR grade H₂O, heat pretreatment equipment (thermoblock, microwave), centrifuge, cold store and thermal block cycler. Buffered solutions for DNA extraction or purification, enzyme treatments, highly sensitive detection systems, and other auxiliary reagents are available from Genova Scientific.

STORAGE AND STABILITY:

Store at -20°C until the expiration date printed on product label. Avoid prolonged exposure to light. Avoid multiple freeze-thaw cycles and exposure to frequent temperature changes. Do not use after the expiration date. If the product is stored under different conditions from those stipulated in these technical indications, the new conditions must be verified by the user. The validity period of the ready to use products when opened, is the same as the expiration date indicated on the label of intact product.

Genova Scientific guarantees that the product will maintain all of the described characteristics from the production date until the expiration date, as long as the product is stored and used as recommended. No other guarantees are provided. Under no circumstances Genova Scientific is obliged to cover damages caused by use of this reagent.



Catalog number



Batch code



Research use only



Temperature limitation



Expiration date



Manufacturer



See instruction for use



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TROUBLESHOOTING:

If unusual amplification is observed or any other deviations from the expected results, please read these instructions carefully, along with the instructions from the PCR system. If this does not solve the problem, please contact Genova Scientific's technical support department (techsupport@genovalab.com) or your local distributor.

PRECAUTIONS:

Use only by qualified personnel.

Use proper protective equipment in order to avoid contact with reagents and samples in the eyes, skin, and mucosal tissues. In case of contact with sensitive areas, immediately flush the affected area with water. Avoid microbial contamination of the reagent, as this may produce nonspecific amplification results.

Material safety data sheet (MSDS) is available upon request.

PERFORMANCE CHARACTERISTICS:

Genova Scientific has performed studies to evaluate the functioning of this polymerase for use with standard amplification systems, concluding that the product is both specific and sensitive for PCR performance.

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