## v*i*vant*i*s DNA AMPLIFICATION PRODUCT



### **Description :**

2X At Tag Master Mix is an optimized ready-to-use 2X concentrated DNA amplification mixture containing At Taq DNA Polymerase, reaction buffer, dNTPs and MgCl<sub>2</sub>. It contains all the components required for routine DNA amplification except template and primers. At Tag DNA polymerase is a complex of specific anti-Taq monoclonal antibody with top quality thermostable Taq DNA Polymerase for automatic "Hot Start" amplification, resulting in greatly improved amplification specificity, sensitivity and yield.

### Features:

- · Saves time and reduces contamination due to reduced number of pipetting steps.
- amplification with enhanced specificity, sensitivity and yield.
- amplification eith reduced artefacts, such as primer-dimer formation and mispriming in multiplex amplification.

### Storage and Stability:

- 2X At Taq Master Mix is stable at -20°C for one year or at 4°C for 18 months if properly stored.
- 2X At Taq Master Mix is stable for 20 freeze-thaw cycles. To avoid frequent freeze-thaw, keeping small aliquot at -20°C is recommended.
- For daily use, keeping an aliquot at 4°C is recommended.

### **Quality Control:**

All preparations are assayed for contaminating endonuclease, exonuclease, and non-specific DNase activities. Functionally tested in DNA amplification.



Amplification of 1.5kb DNA fragment from pTZ DNA region using 2X At Taq Master Mix in a 50µl reaction mixture.

- Lane M : VC 1kb DNA Ladder.
- Lane 1 : DNA amplification product generated with 1.25u of At Taq DNA polymerase.
- Lane 2 : DNA amplification product generated with
  - 2X At Tag Master Mix (store at -20°C).
- Lane 3 : DNA amplification product generated with 2X At Taq Master Mix (after 20 freeze-thaw cycles).

1.0% TBE agarose gel.



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Add the following components in a 0.2ml thin walled PCR tube on ice. Gently mix all solutions after thawing. Spin down briefly and keep on ice X

## For 50µl reaction volume:

later, nuclease-free	DNA Template	rimers (Fwd / Rev)	MgCl <sub>2</sub> (50mM)	2X AT Taq Master Mix	Reagent:	
Adjust final vc	Variable	Variable	Refer to Table (A)	25µl	Volume	
lume to 50µl	0.02 - 5µg	0.1 - 1 µM each	**For more than 1.5mM MgCl <sub>2</sub>	*1X	Final Concentration	

Note

: Smaller reaction

Extension / 1kb Final Extension

Annealing

50 - 68°C for 30 seconds

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- 35 cycles

94°C for 30 seconds 94°C for 2 minutes

72°C for 30 seconds

72°C for 7 minutes

This protocol may change depending on the template DNA and primers used

Denaturation

Denaturation

\*2X At Taq Master Mi

adding additional MgCl<sub>2</sub>. Please refer to Table (A) if higher MgCl<sub>2</sub> concentration is preferred

ration of 1.5mM. Howe

higher concentration may be achieved

g

2.5

4.0

volume may be achieved provided that the same final concentration of each reaction component is maintained

# Table (A) : For more than 1.5mM final MgCl<sub>2</sub> concentration

2.0	1.5	1.0	0.5	Volume of MgCl_2 (50mM) stock to add into 50 $\mu l$ reaction mixture ( $\mu l$ )
3.5	3.0	2.5	2.0	Final $MgCl_2$ concentration (mM)

Product Use Limitation This product is for research purpose an in vitro use only v*i*vant*i* s

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