

2X ViRed Taq Master Mix

Product No : CLMM01
Quantity : 100 reactions



Lot :
Expiry Date :
Supplied with : 4 x 625µl 2X ViRed Taq Master Mix*
3ml of Nuclease-free Water
1ml of 50mM MgCl₂

Store at -20°C
*2X ViRed Taq Master Mix consists of Taq DNA Polymerase, Vibuffer A, dNTPs, MgCl₂, inert red dye and stabilizers.



info@vivantechnologies.com

Description :

2X ViRed Taq Master Mix is an optimized ready-to-use 2X concentrated DNA amplification mixture premixed with red color tracking dye. The ViRed Taq Master Mix contains Taq DNA Polymerase, reaction buffer, dNTPs, MgCl₂, inert red dye and stabilizers needed for routine DNA amplification to obtain a wide range of PCR and DNA products up to 8kb. An inert red dye and stabilizers allows direct loading of final products onto gels for electrophoresis. The red color dye migrates at approximately 400bp on 1% agarose in 1X TBE Buffer.

Features:

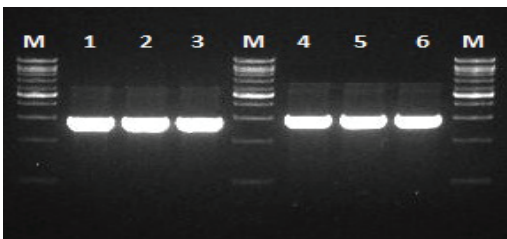
- Suitable for all routine DNA amplification applications
- Reduces set-up time and buffer-dye mixing
- Minimizes potential contamination by eliminating several pipetting steps
- Easy confirmation of complete mixing
- No additional loading dye needed – direct loading of final products onto gels

Storage and Stability:

- Stable at -20°C for 18 months or at 4°C for 6 months if properly stored
- Stable for 20 freeze-thaw cycles. To avoid frequent freeze-thaw, keeping small aliquots at -20°C is recommended
- For daily use, keeping aliquots 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 ViRed Taq Master Mix in a 50µl reaction mixture (1.0% TBE agarose gel).

Lane M : VC 1kb DNA Ladder
Lane 1 : DNA amplification product generated with 1.25u of Taq DNA Polymerase
Lane 2 : DNA amplification product generated with 2X ViRed Taq Master Mix (store at -20°C)
Lane 3 : DNA amplification product generated with 2X ViRed Taq Master Mix (after 20 freeze-thaw cycles)
Lane M : VC 1kb DNA Ladder
Lane 4 : DNA amplification product generated with 1.25u of Taq DNA Polymerase
Lane 5 : DNA amplification product generated with 2X Taq Master Mix (store at -20°C)
Lane 6 : DNA amplification product generated with 2X Taq Master Mix (after 20 freeze-thaw cycles)

RECOMMENDED PROTOCOL FOR 2X ViRed Taq Master Mix:
Gently mix all solutions after thawing. Spin down briefly and keep on ice. Add the following components in a 0.2ml thin walled PCR tube on ice.
For 50µl reaction volume:

Reagent:	Volume	Final Concentration
2X ViRed Taq Master Mix	25µl	*1X
MgCl ₂ (50mM)	Refer to Table (A)	**For more than 1.5mM MgCl ₂
Primers (Fwd / Rev)	Variable	0.1 - 1 µM each
DNA Template	Variable	0.02 - 5µg
Water, nuclease-free	Adjust final volume to 50µl	

**2X ViRed Taq Master Mix contains a fixed final MgCl₂ concentration of 1.5mM. However, higher concentration may be achieved by adding additional MgCl₂. Please refer to Table (A) if higher MgCl₂ concentration is preferred.
Note : Smaller reaction volume may be achieved provided that the same final concentration of each reaction component is maintained.

CYCLING CONDITIONS (100bp-5kb)	
Denaturation	94°C for 2 minutes
Denaturation	94°C for 30 seconds
Annealing	50 - 68°C for 30 seconds
Extension / 1kb	72°C for 30 seconds
Final Extension	72°C for 7 minutes

} 25 - 35 cycles
This protocol may change depending on the template DNA and primers used.

Table (A) : For more than 1.5mM final MgCl₂ concentration

Volume of MgCl ₂ (50mM) stock to add into 50µl reaction mixture (µl)	Final MgCl ₂ concentration (mM)
0.5	2.0
1.0	2.5
1.5	3.0
2.0	3.5
2.5	4.0