

Suggested PCR Protocol

Pipette the following into a PCR tube, mix and make up to a final volume of 50µL. We recommend dispensing all reagents on ice, adding the enzyme last. It is important to vortex all buffers and MgCl₂ solutions before use to remove any gradients that may result from repeated freeze/thaw steps. If you do have already your own PCR-Protocol established, please use your existing pipetting scheme and Thermocycler protocol.

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Components	Quantities
Template DNA	1 ng - 10 ng plasmid DNA or 5 ng - 500 ng genomic DNA
Nucleotides	1µL (10 mM) each dNTP
10X amplification buffer	5µL
primer 1:	4-7µL of 3 µM solution (10 - 20 pmol absolute)
primer 2:	4-7µL of 3 µM solution (10 - 20 pmol absolute)
sterile, bidistilled water	up to 50µL
<i>Pwo</i> -Polymerase	0.3 - 0.8µL (0.5 - 2.0 units)

The following cycling protocol has to be seen as suggestion. Cycle numbers and cycle times have to be adjusted.

Cycling Profile *	
94°C for 2-4 minutes	step 1(denaturing of DNA)
	(cycling part)
94°C for 30-60 sec.	1 - 30 cycle(s)
annealing temp. for 30-60 sec.	1 - 30 cycle(s)
extension (72°C) for 60-240 sec. **	1 - 30 cycle(s)
	final step (filling up step)
94°C for 30 sec.	
72°C for 5-10 minutes	
cool down to 4°C	

* cycling times are proposals and have to be adjusted to the special needs.

** the longer the template to be amplified, the longer the amplification time

After running the amplification reaction, 5 µL of each sample should be analyzed in a 1% agarose gel containing 0.5µg/mL of ethidium bromide.

Note: For every template/primer pair the optimal reaction conditions have to be evaluated empirically, changing the primer/template ratio, the ionic strength (with MgSO₄) and the cycle parameters (time and temperatures).

Trouble shooting:

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Check if annealing temperature has been calculated correctly. The optimal annealing temperature of primers lies about 5°C below the T_M of the oligo.

TM calculation: For oligos up to 15 bases: $TM = 4x(G+C) + 2x(A+T)$
 For larger oligos: $TM = 81,5+16,6([\log_{10}J^*]) + 0,4(\%G+C) - (600/n) - 0,63(\%FA)$

A = Adenosine, C = Cytosine, G = Guanine, T = Thymidine, FA = Formamide, J* = concentration of monovalent cations, n = number of bases; T_M = calculated melting temperature,

Remark: If primers with mismatches are used the *Pwo* Polymerase must be added immediately before the start of the PCR®. Do not let the reaction mix stand at room temperature since primers will be degraded by the *Pwo* Polymerase.

Bands smear over the Gel

- Increase annealing temperature
- Add up to 5% DMSO to the Reaction mixture
- Reduce the amount of template DNA
- Check if the primers bind more than once on the template DNA. Perform control reactions with only one of the primers.

Low Yields

- Increase annealing temperature

The PCR process is licensed and covered under US-Patent owned by Hoffmann-La Roche, Roche Int. USA

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