

Getting the Most Out of Your PCR

A best practice and optimization guide



Your PCR workflow ...

- 1 DNA concentration**
Measure your DNA (gDNA/ cDNA) for concentration and purity.
Ideal DNA purity range (A_{260}/A_{280}) = ~1.7 – 2.0
 - 2 Calculate concentration**
Calculate according to desired reaction volume.
 - 3 Mastermix**
Prepare a mastermix of all common components in 1 tube. Remember to always use suitable controls (positive, negative, no template, etc.).
 - 4 Mix**
Mix well to ensure homogeneous distribution of components to minimize well-to-well variation.
(See Eppendorf Application Note 130 for best mixing guide.)
 - 5 Briefly spin down (short-spin)**
Short spin can reduce contamination by pulling down the liquid from the rim of the tube.
 - 6 Distribute mastermix**
Hold pipette vertically during pipetting. Always dispense to the bottom of the tube.
 - 7**
- > Set cycler modes to match reaction volume (fast, standard, safe) for optimal heat transfer
> Denaturation: temperature depends on enzyme; holding time depends on enzyme, DNA source and GC content
> Annealing: temperature generally ~5-10°C below primer melting temperature. Long holding time not necessary.
> Extension: optimal temperature for polymerase activity (e.g. Taq = 72°C). Incorporation rate of Taq: ~1000 bases/ min.

... and tips & tricks for some steps

- 1**
> Take note of the absorbance reading (not just concentration values). Aim for 0.1 - 1.0 A for reliable reading according to the Beer-Lambert Law.
- 2**
> Aliquot stock solutions to prevent multiple freeze-thawing events that can affect reproducibility of PCR.
> Smaller reaction volume saves reagents and costs, but when working in small volumes it is important to protect against evaporation during PCR.
> Use low retention tips or low binding tubes to minimize sample and reagent loss.
- 3**
> Prepare each mastermix in a single tube to prevent pipetting variation
> Use bigger tube sizes when preparing mastermixes:
(e.g. 1 x 5 mL tube is better than 2 x 2 mL tubes).
- 4**

Use PCR clean consumables that are certified:
> Human DNA-free > DNase-free
> RNase-free > Free of PCR inhibitors

- 5**
> Use a dedicated set of pipettes for PCR.
> Use the appropriate tips for the pipettes.
> Calibrate the pipettes at least once a year to ensure accuracy and precision.
> Prevent aerosol contamination by using either positive displacement pipette or filtered tips.
- 7**
Always take note of ramp rates when:
> Transferring protocols between different cyclers.
> Transferring from optimization protocol to standard protocol.
> Set cycler temperature modes to match reaction volume (e.g. safe, fast, standard in Eppendorf cyclers).

Troubleshooting your PCR

Problems	Possible solutions
1. Non-specific amplifications	<ol style="list-style-type: none"> 1. Use Hot-start strategies: <ol style="list-style-type: none"> a) Manual hot-start b) Use devices with thermal sample protection (TSP) lid c) Use devices with "Impulse PCR" function d) Use hot-start reagents 2. For new primers, run optimization with single-primer (e.g. forward primer only) controls to determine non-specificity 3. Alternative strategies: <ol style="list-style-type: none"> a) Mg2+ titration (concentration optimization) b) Touchdown PCR
2. No / low amplifications	<ol style="list-style-type: none"> 1. Optimize denaturation and/or annealing temperature with gradient function 2. Use PCR enhancers (e.g. DMSO, BSA). These require empirical testing for each combination of template and primer